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5.1.6
Compugen Ltd.
GenCore version (c) 1993 - 2005
           Copyright
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- protein search, using sw model OM protein

September 21, 2005, 13:41:21; Search time 39 Seconds (without alignments) 268.914 Million cell updates/sec Run on:

US-10-700-314-2 109 Title: Perfect score:

1 MKFISTSLLLMLLVSSLSPV......VLRKRSSSTLPVPVFKRKIP 109 Sequence:

Scoring table:

OLIGO . Gapop 60.0 , Gapext 60.0

283416 segs, 96216763 residues Searched:

9 Word size :

Total number of hits satisfying chosen parameters:

529

Minimum DB seq length: 0 Maximum DB seq length: 2000000000

Post-processing: Listing first 45 summaries

Database :

PIR 79:*
1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

•		de			SUMMARIES	
Result No.	Score	Query Match	Length	DB	ID	Description
1	6	8.3	1139	2	A49370	ElA-associated cyc
	80	7.3	102	7	T36017	probable SecG-like
м	7	6.4	90	7	F82571	hypothetical prote
4	7	6.4	94	7	T03751	_
S	7	6.4	174	7	T08702	
9	7	6.4	235	N	A99603	hypothetical prote
. 7	7	6.4	312	~	H70107	methionyl-tRNA for
œ	7	6.4	331	~	AE1118	dihydroxyacetone k
თ	7	6.4	331	7	AG1478	dihydroxyacetone k
10	7	6.4	398	~	A45633	rhoptry-associated
11	7	6.4	427	7	F64084	3-deoxy-manno-octu
12	7	6.4	496	7	C64935	hypothetical prote
13	7	6.4	496	7	E90936	
14	7	6.4	496	7	A85785	
15	7	6.4	521	7	E95052	hypothetical prote
16	7	6.4	570	7	S70400	ъ
17	7	6.4	608	7	B70188	н
18	7	6.4	722	7	S64492	
19	7	6.4	738	7	E95952	д
20	7	6.4	837	7	T19271	hypothetical prote
21	7	6.4	958	7	T44046	
22	7	6.4	1227	~	A86245	_
23	7	6.4	1378	Н	148751	protein-tyrosine k
24	7	6.4	2948	N	T22664	hypothetical prote
25	9	5.5	40	7	H95063	hetical
26	9	5.5	57	-	SHNZE2	small hydrophobic
27	9	5.5	57	Н	SHNZE4	small hydrophobic
28	9	5.5	65	7	S08014	calcium-binding pr
	9	5.5	69	7	G82496	O

transforming growt hypothetical prote NADH2 dehydrogenas	<pre>sperm-specific pro peptide YY precurs conserved hypothet</pre>	very hypothetical sperm-specific pro	hypothetical prote hypothetical prote conserved hypothet	hypothetical prote hypothetical prote	hypothetical proce mating pheromone a
A40699 T22936 T11163	S34115 A55914 H81710	T38892 S65494	G71202 H71209 E87535	T12461 S75884	B/2633 D97328 S05791
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71 83 91	92	101	104 107 109	110	119
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30 31 32	33 34 54	36	338 4 3 9 8 0	4 4 4 1 2 4	4 4 4 5 4 5

ALIGNMENTS

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RESULT 1
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ElA-associated cyclin-binding protein p130 - human N/Alternate names: adenovirus ElA-associated 130K protein; retinoblastoma-like protein 2

C;Species: Nome sapiens (man)
C;Species: Home sapiens (man)
C;Species: Home sapiens (man)
C;Species: Home sapiens (man)
C;Date: 07-Apr-1994 #sequence revision 18-Nov-1994 #text_change 09-Jul-2004
C;Accession: A49370; A49369; I38150; S36648
R;Hannon, G.J.; Demetrick, D.; Beach, D.
A;Hannon, G.J.; Demetrick, D.; Beach, D.
A;Title: Isolation of the Rb-related pl30 through its interaction with CDK2 and cyclins.
A;Reference number: A49370; MUID:94074896; PMID:8253384

A;Status: proliminary
A;Rasidues: 1-1139 <HANA
A;Residues: 1-1139 <HANA
A;Residues: 1-1139 <HANA
A;Residues: 1-1139 <HANA
A;Residues: UNIPROT:Q08999; GB:S67171; NID:g453131; PIDN:AAB29227.1; PID:g453132
A;Cross-references: UNIPROT:Q08999; GB:S67171; NID:g453131; PIDN:AAB29227.1; PID:g453132
A;Experimental source: Hela S3 suspension cells
A;Experimental source: Hela C3 suspension cells
A;Note: sequence extracted from NCBI backbone (NCBIN:140577)
B;Li, Y: Graham, C.; Lacy, S.; Duncan, A.M.V.; Whyte, P.
A;Licle: The adenovirus EIA-associated 130-kD protein is encoded by a member of the reti
A;Reference number: A49369; MUID:94074895; PMID:8253383

A;Status: preliminary

A; Molecule type: mRNA A; Residues: 1-36,'S',38-63,'P',65-658,'RL',661-1139 <LIA> A; Cross-references: GB:X76061 R; MayOl, X.; Grana, X.; Baldi, A.; Sang, N.; Hu, Q.; Giordano, A. Oncogene 8, 2561-2566, 1993 A; Title: Cloning of a new member of the retinoblastoma gene family (pRb2) which binds to A; Reference number: 138150; MUID:93368960; PMID:8361765

A;Accession: I38150
A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule (rype: mRNA
A;Residues: 58-205,'M', 207-1139 <MAY>
A;Residues: 58-205,'M', 207-1139 <MAY>
A;Ccoss-references: EMBL:X74594; NID:g397147; PIDN:CAA52671.1; PID:g397148

A;Cross-references: GDB:269890; OMIM:180203 A;Map position: 16q12.2-16q12.2 C;Superfamily: retinoblastoma-associated protein C;Keywords: cell cycle control

Gaps ö Length 1139; 0; Indels 8.3%; Score 9; DB 2; 100.0%; Pred. No. 0.55; tive 0; Mismatches Query Match 8.3 Best Local Similarity 100. Matches 9; Conservative

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94 RSSSTLPVP 102 970 RSSSTLPVP 978 셤 ð

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C;Species: mitochondrion Physarum polycephalum
C;Date: 23-Apr-1999 #sequence_revision 23-Apr-1999 #text_change 09-Jul-2004
C;Accession: T03751
R;Nakagawa, C.C.; Jones, E.P.; Miller, D.L.
Curr. Genet. 33, 178-187, 1998
A;Title: Mitochondrial DNA rearrangements associated with mF plasmid integration and plasmite number: Z15055; MUID:98177147; PMID:9508792
                                                                                                                                                                                                                                                                                                     A;Cross-references: UNIPROT:O20627; EMBL:AF012249; NID:g2435419; PIDN:AAC15931.1; PID:g24.
A;Experimental source: strain aux2-S
C;Genetics:
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Cincoession: A95004
Cincoession: A95004
Cincoession: A95003
Cincoession: A95003
A; Title: The complete genome sequence of the murine respiratory pathogen Mycoplasma pulmc A; A; Cereanion: A99603
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A;Experimental source: strain UAB CTIP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       hypothetical protein MYPU 7290 [imported] - Mycoplasma pulmonis (strain UAB CTIP)
C;Species: Mycoplasma pulmonis
C;Date: 24-May-2001 #sequence_revision 24-May-2001 #text_change 09-Jul-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   C;Species: Homo sapiens (man)
C;Date: 11-Jun-1999 #sequence_revision 11-Jun-1999 #text_change 09-Jul-2004
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C; Accession: T08702

Swanbutt. R.; Heubner, D.; Mewes, H.W.; Gassenhuber, J.; Wiemann, submitted to the Protein Sequence Database, March 1999

A; Reference number: 216471

A; A; Accession: T08702

A; Molecule type: MRNA

A; Residues: 1-174 < WAM>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       6.4%; Score 7; DB 2; Length 235;
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A,Experimental source: fetal brain; clone DKFZp5640243
C,Genetics:
A,Note: DKFZp5640243.1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    hypothetical protein DKFZp5640243.1 - human (fragment)
                                                                                                                                                                                                                                           A;Accession: T03751
A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
A;Residues: 1-94 <NAK>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 2;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      0; Mismatches
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100.0%; Pred. No.
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Best Local Similarity 100.
Matches 7; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity 100.
Matches 7; Conservative
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A; Residues: 1-235 < KUR>
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A;Genetic code: SGC3
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A,Experimental source: strain 935c
R,Simpson, A.JG; Reinach, F.C.; Arruda, P.; Abreu, F.A.; Acencio, M.; Alvarenga, R.; Briones, M.R.S.; Bueno, M.R.P.; Camargo, A.A.; Camargo, L.R.; Carraro, D.M.; Carrer, H. assonet, C.; El-Dorry, H.; Facincani, A.P.; Ferreira, A.J.S.
abovero, E.; Docena, C.; El-Dorry, H.; Fracincani, A.P.; Franca, S.C.; Franco, D.M.; Carrer, H. abunitted to GenBank, June 2000, M.S.; Franca, S.C.; Franco, M.C.; Frohm J.D.; Junqueira, M.L.; Kemper, E.L.; Kitajima, J.P.; Krieger, J.E.; Kuramae, E.E.; Laigr J.D.; Junqueira, M.A.; Madeira, A.M.B.N.; Matcha, M.A.; Madeira, A.M.B.N.; Matcha, M.A.; Madeira, A.M.B.N.; Matcha, M.A.; Matcha, M.A.; Matcha, B.M.F.; Matukuma, A.Y.; Menck, C.F.M.; Miyaki, C.Y.; F.G.; Nunes, L.R.; Oliveira, M.A.; de Oliveira, M.C.; de Oliveira, R.C.; Palmieri, D.A. Rofigues, V.; Rosa, A.J.; de Silva, A.M.; Silva, A.M.; Salvasak, A.A. Tsuhako, M.H.; Vallada, H.; Van Sluys, M.A.; Verjovski-Almeida, S.; Vettore, A.L.; Z.
                                                                                                                                                                                                                                                                                                                                                                                                 A;Residues: 1-102 <SEE>
A;Residues: 1-102 <SEE>
A;Cross-references: UNIPROT:Q9Z521; EMBL:AL035591; PIDN:CAB38134.1; GSPDB:GN00070; SCOED & Experimental source: strain A3(2)
C;Genetics: A;Gene: SCOEDB:SCC54.04c
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               hypothetical protein XP2321 [imported] - Xylella fastidiosa (strain 9a5c)
C;Species: Xylella fastidiosa
C;Species: Xylella fastidiosa
C;Date: 18-Aug-2000 #sequence_revision 20-Aug-2000 #text_change 09-Jul-2004
C;Accession: F82571
R;anonymous, The Xylella fastidiosa Consortium of the Organization for Nucleotide Sequen Nature 406, 151-157, 2000
A;Title: The genome sequence of the plant pathogen Xylella fastidiosa.
A;Reference number: A82515; MUID:20365717; PMID:10910347
A;Note: for a complete list of authors see reference number A59328 below
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hypothetical protein A - slime mold (Physarum polycephalum) mitochondrion integrated pla
                                                                    probable SecG-like integral membrane protein - Streptomyces coelicolor C; Species: Streptomyces coelicolor C; Species: Streptomyces coelicolor C; Date: 03-Dec-1999 #sequence_revision 03-Dec-1999 #text_change 09-Jul-2004 C; Accession: T36017 R; Seeger, K.; Harris, D.; Bentley, S.D.; Parkhill, J.; Barrell, B.G.; Rajandream, M.A.submitted to the EMBL Data Library, March 1999
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0
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0
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                                                                                                                                                                                                                                                                      A;Reference number: 221581
A;Accession: T36017
A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   7.3%; Score 8; DB 2;
Local Similarity 100.0%; Pred. No. 0.76;
Les 8; Conservative 0; Mismatches
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100.0%; Pred. No. 7.7;
iive 0; Mismatches
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Best Local Similarity
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A;Molecule type: DNA
A;Residues: 1-90 <SIM>
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3 FISTSLL 9

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Best Local Similarity Matches 7; Conserv

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C;Species: Listeria innocua Linuage Lampoiteu; - Listeria Linuacua (Strain Cilpli262)
C;Species: Listeria innocua
C;Date: 27-Nov-2001 #sequence_revision 27-Nov-2001 #text_change 09-Jul-2004
C;Accession: AG1478
C;Accession: AG1478
C;Baser, P:, Frangeul, L.; Buchrieser, C.; Amend, A.; Baquero, F.; Berche, P.; Bloecker.; Dominguez-Bernal, G; Durand, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H. D.; Jones, L.M.; Karst, U.
Science 294, 849-852, 2001
A;Authors: Kreft, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.; Ma ok, C.; Schlueter, T.; Simoes, N.; Tierrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehland, A;Reference number: Allo77; MUID:21537279; PMID:11679669
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            A;Cross-references: UNIPROT:Q92EU2; GB:AL592022; PIDN:CAC95599.1; PID:g16412795; GSPDB:G
A;Experimental source: strain Clip11262
C;Genetics:
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C;Species: Plasmodium falciparum
C;Species: Plasmodium falciparum
C;Date: 22-Apr-1993 #sequence_revision 18-Nov-1994 #text_change 09-Jul-2004
C;Accession: A45633
R;Saul, A.; Cooper, J.; Hauquitz, D.; Irving, D.; Cheng, Q.; Stowers, A.; Limpaiboon, T. Mh. Biochem. Parasitol. 50, 139-149, 1992
A;Title: The 42-kilodalton rhoptry-associated protein of Plasmodium falciparum.
A;Reference number: A45633; MUID:92178281; PMID:1542308
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       dihydroxyacetone kinase homolog lin0366 [imported] - Listeria innocua (strain Clip11262)
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A;Authors: Gnehm, C.L.; McDonald, L.A.; Small, K.V.; Fraser, C.M.; Smith, H.O.; Venter, A;Title: Whole-genome random sequencing and assembly of Haemophilus influenzae Rd.
A;Reference number: A64000; MUID:95350630; PMID:7542800
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C;Date: 18-Aug-1995 #sequence_revision 18-Aug-1995 #text_change 09-Jul-2004
C;Accession: F64084
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A;Molecule type: DNA; protein
A;Residues: 1-39 cSAU3
A;Cross-references: UNIPROT:Q26004
A;Cross-reference extracted from NCBI backbone (NCBIN:85242, NCBIP:85246)
C;Superfamily: Plasmodium falciparum rhoptry-associated protein 2
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100.0%; Pred. No. 24;
ve 0; Mismatches
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100.0%; Pred. No. 28;
tive 0; Mismatches
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Best Local Similarity 100.
Matches 7; Conservative
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Matches 7; Conservative
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A;Molecule type: DNA
A;Residues: 1-331 <GLA>
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dihydroxyacetone kinase homolog lmo0348 [imported] - Listeria monocytogenes (strain EGD-C; Species: Listeria monocytogenes
C; Species: Listeria monocytogenes
C; Date: 27-Nov-2001 #text_change 09-Jul-2004
C; Accession: AE1118
R;Glaser, P.; Frangeul, L.; Buchrieser, C.; Amend, A.; Baquero, F.; Berche, P.; Bloecker, F.; Pominguez-Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H. D.; Jones, L.M.; Karst, U.
Science 294, 849-882, 2001
A; Authors: Kreft, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.; Maok, C.; Schlueter, T.; Simoes, N.; Tierrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehland, A; Reference number: AB1077; MUD:21537279; PMID:11679669
A; Rocession: AE1118
A; Steaus: preliminary
A; Molecule type: DNA
A; Residues: 1-331 cGLA>
A; Cross-references: UNIPROT:Q8YA18; GB:NC_003210; PIDN:CAC98427.1; PID:g16409726; GSPDB: A; Rocessioner: Imo0348
                                                                                                                                                                                                                                                                                                                                                                   C;Species: Borrelia burgdorferi (Lyme disease spirochete)
C;Species: Borrelia burgdorferi (Lyme disease spirochete)
C;Dectes: 13-Feb-1998 #sequence_revision 13-Feb-1998 #text_change 09-Jul-2004
C;Accession: H70107
R;Fraser, C.M.; Casjens, S.; Huang, W.M.; Sutton, G.G.; Clayton, R.; Lathigra, R.; White son, D.; Peterson, J.; Kerlavage, A.R.; Quackenbush, J.; Salzberg, S.; Hanson, M.; Vugt, Bown, C.; Garland, S.; Pujii, C.; Cotton, M.D.; Horst, K.; Roberts, K.; Hatch, B. Nature 390, 580-586, 1997
A;Authors: Smith, H.O.; Venter, J.C.
A;Title: Genomic sequence of a Lyme disease spirochaete, Borrelia burgdorferi.
A;Reference number: A70100; MUID:98065943; PMID:9403685
A;Accession: H70107
A;Reference number: A70100; MUID:98065943; PMID:9403685
A;Accession: H70107
A;Reference number: A70100; MUID:98065943; PMID:9403685
A;Accession: H70107
A;Rolecule type: DNA
A;Residues: 1-312 <KLE>
A;Cross-references: UNIRROF:051091; GB:AE001119; GB:AE000783; NID:g2687936; PIDN:AAC6644
A;Experimental source: strain B31
C;Superfamily: methionyl-tRNA formyltransferase; phosphoribosylglycinamide formyltransfe
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H70107
methionyl-tRNA formyltransferase (EC 2.1.2.9) - Lyme disease spirochete
C;Species: Borrelia burgdorferi (Lyme disease spirochete)
C;Date: 13-Feb-1998 #sequence_revision 13-Feb-1998 #text_change 09-Jul-2004
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                                       Indels
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100.0%; Pred. No. 24;
iive 0; Mismatches
                                          0; Mismatches
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Best Local Similarity 100.0
Matches 7; Conservative
                                       7; Conservative
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165 LVSSLSP 171

13 LVSSLSP 19

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Local Similarity hes 7; Conserv

Query Match Best Loc Matches SSSTLPV 101 SSSTLPV 198

95 192

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probable transport system permease protein Z2787 [imported] - Escherichia coli (strain O) C;Species: Escherichia coli (parei 16-Feb-2001 #sequence_revision 16-Feb-2001 #text_change 09-Jul-2004 C;Accession: A85785 R;Perna, N.T.; Plunkett III, G.; Burland, V.; Mau, B.; Glasner, J.D.; Rose, D.J.; Mayhew, iller, L.; Grotbeck, E.J.; Davis, N.W.; Lim, A.; Dimalanta, E.; Potamousis, K.; Apodaca, A;Title: Genome sequence of enterohemorrhagic Escherichia coli O157:H7.
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C;Genetics:
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Gipeciae: Streptococcus pneumoniae
Gipeciae: Streptococcus pneumoniae
Gipeciae: Streptococcus pneumoniae
Gipaciae: Streptococcus pneumoniae
Gipacession: B9502
Rifettelin, H.; Nelson, K.E.; Paulsen, I.T.; Eisen, J.A.; Read, T.D.; Peterson, S.; Heide on, J.D.; Umayam, L.A.; White, O.; Salzberg, S.L.; Lewis, M.R.; Radune, D.; Holtzapple, I nson, T.; Hickey, B.K.; Holt, I.E.
Science 293, 498-506, 2001
A, Authors: Loftus, B.J.; Yang, F.; Smith, H.O.; Venter, J.C.; Dougherty, B.A.; Morrison, A; Title: Complete Genome Sequence of a virulent isolate of Streptococcus pneumoniae.
A, Reference number: A95000; MUID:21357209; PMID:11463916
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Best Local Similarity 100...
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A, Status: preliminary
A, Molecule type: DNA
A, Residues: 1-496 <STO>
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A;Molecule type: DNA
A;Residues: 1-521 <KUR>
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C64335
Cypochetical protein b1755 - Escherichia coli (strain K-12)
Cypochetical protein b1755 - Escherichia coli (strain K-12)
Cypochetical protein a coli
Cypochetical protein a coli
Cypochetical Sep-1997 #sequence_revision 17-Sep-1997 #text_change 01-Mar-2002
CyAccession: C64335
R:Blattner, F.R.; plunkett III, G.; Bloch, C.A.; Perna, N.T.; Burland, V.; Riley, M.; Cc. A.; Rose, D.J.; Mau, B.; Shao, Y.
Science 277, 1453-1462, 1997
A; Ttle: The complete genome sequence of Escherichia coli K-12.
A; Reference number: A64720; MUID:97426617; PMID:9278503
A; Accession: C64935
A; Accession: C64935
A; Accession: C64935
A; Catuus: preliminary; nucleic acid sequence not shown; translation not shown
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A;Experimental source: strain Ol57:H7, substrain RIMD 0509952
C;Genetics:
A;Gene: EC82461
A;Residues: 1-427 <TIGR>
A;Cross-references: UNIPROT:P44806; GB:U32748; GB:L42023; NID:g1573645; PIDN:AAC22311.1; C;Function:
C;Function: catalyzes the transfer of 3-deoxy-D-manno-octulosonic acid from its CMP c;Superfamily: Chlamydia trachomatis 3-deoxy-manno-octulosonate cytidylyltransferase C;Keywords: nucleotidyltransferase; transmembrane protein
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X;Residues: 1-496 <BLAT>
A;Cross-references: GB:AE000270; GB:U00096; NID:g1788045; PIDN:AAC74825.1; PID:g1788052;
A;Experimental source: strain K-12, substrain MG1655
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100.0%; Pred. No. 34;
tive 0; Mismatches
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100.0%; Pred. No. 34;
tive 0; Mismatches
                                                                                                                                                                                                                    Query Match 6.4%; Score 7; DB 2; Best Local Similarity 100.0%; Pred. No. 30; Matches 7; Conservative 0; Mismatches
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Best Local Similarity 100.º
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Matches 7; Conservative
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Gaps

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0; Indels

Length 496;

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A;Cross-references: UNIPROT:Q97SD1; GB:AE005672; PIDN:AAK74614.1; PID:g14971923; GSPDB:GA
A;Experimental source: strain TIGR4
C;Genetics:
A;Gene: SP0453
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                                                                                                                                    Gaps
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0
                                                                                                 Length 521;
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                                                                                                     DB 2;
                                                                                                 6.4%; Score 7; DB 2;
100.0%; Pred. No. 35;
:ive 0; Mismatches
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RESULT 14

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September 21, 2005, 13:41:20 ; Search time 175 Seconds (without alignments) 318.952 Million cell updates/sec
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1 MKFISTSLLLMLLVSSLSPV......VLRKRSSSTLPVPVFKRKIP 109
GenCore version 5.1.6 (c) 1993 - 2005 Compugen Ltd.
                                                                              OM protein - protein search, using sw model
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                   Copyright
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                                                                                                                    Run on:
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OLIGO Gapop 60.0 , Gapext 60.0

Scoring table:

1612378 seqs, 512079187 residues Searched:

9

Word size :

Total number of hits satisfying chosen parameters: Minimum DB seq length: 0 Maximum DB seq length: 200000000

2944

Post-processing: Listing first 45 summaries

1: uniprot sprot:*
2: uniprot_trembl:* UniProt 03:* Database :

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

	Description	043927 homo sapien	Q8hyn9 macaca mula	Q7ys87 macaca mula	Q7z312 homo sapien			Q9z521 streptomyce						Q8fk19 escherichia						helicoba			Q8qz54 sabo virus.	20627 physarum po		Q86cz9 caenorhabdi				Q8p8j3 xanthomonas	enterococ	Q8m172 tortanus fo
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de	Query Match	100.0	21.1	17.4	8.3	8.3	8.3	7.3	7.3	7.3	7.3	7.3	7.3	7.3	7.3	7.3	7.3	7.3	6.4	6.4	6.4	6.4	6.4	6.4	6.4	6.4	6.4	6.4	6.4	6.4	6.4	6.4
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Q9nyd2 homo sapien Q8nu2 methanosarc Q972x9 sulfolobus G7xa30 tortanus fo Q733j8 bacillus ce Q9t6j3 lepus comus Q9t6j3 mycoplasma Q9t6i6 lepus oiost Q6uxu8 homo sapien Q85ju disogmus ar Q9puy2 apteronotus Q9vc67 drosophilus Q7vkq1 haemophilus ce Q732h2 bacillus ce
Q9NYD2 Q9TMU2 Q972X9 Q7XA30 Q7X3J48 Q9T616 Q9T616 Q6UXU8 Q85JU4 Q85JU4 Q9VC67 Q7VKQ1
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## ALIGNMENTS

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TISSUESKeletal muscle;

TISSUESKeletal muscle;

MEDINEL-238825; PubMed=12477932; DOI=10.1073/pnas.242603899;

MEDINEL-238825; PubMed=12477932; DOI=10.1073/pnas.242603899;

MA Strausherg R.D., Feingold E.A., Grouse L.H., Derge J.G.,

Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,

Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,

A plachenko L., Marusina K., Farmer A.A., Rubin G.M., Heigh F.,

A patchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,

A stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,

A stapleton M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,

A Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,

A Rohards S., Worley D.M., McKernan K.J., Malek J.A., Gunaratne P.H.,

A Richards S., Worley C.M., Sodergren E.J., Lu X., Gibbs R.A.,

A Richards S., Worley M.M., Sodergren E.J., Lu X., Gibbs R.A.,

A Hilalon D.K., Murny D.M., Madan A., Rodrigues S., Sanchez A.,

Rah Fahey J., Helton E., Ketteman M., Madan A., Rodrigues S., Sanchez A.,

Mhiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SEQUENCE FROM N.A.
TISSUE-Peripheral blood leukocytes;
MEDLINE-Peripheral blood leukocytes;
MEDLINE-Peripheral blood leukocytes;
Legler D.F., Loetscher M., Stuber Roos R., Clark-Lewis I.,
Baggiolini M., Moser B.;
Baggiolini M., Moser B.;
Bagliolini M., and the separate and the 
                                                                                                                                      30-MAY-2000 (Rel. 39, Created)
30-MAY-2000 (Rel. 39, Last sequence update)
05-MAY-2000 (Rel. 44, Last annotation update)
Small inducible cytokine B13 precursor (CXCL13) (B lymphocyte chemoattractant) (CXC chemokine BLC) (B cell-attracting chemokine 1) (BCA-1) (ANGIE).
                                                                                                                                                                                                                                                                                                                                                                    Name-CXCLIJ; Synonyms=BCAl, BLC, SCYBL3;
Homo sapiens (Human).
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    "A B-cell-homing chemokine made in lymphoid follicles activates
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      SEQUENCE FROM N.A.

BEDILINE=98146056; PubMed=9486651; DOI=10.1038/35876;
Gunn M.D., Ngo V.N., Ansel K.M., Ekland E.H., Cyster J.G.,
Williams L.T.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SEQUENCE FROM N.A.
TISSUE=Peripheral blood;
Napolitano M., Spinetti G., Gaetano C., Capogrossi C.M.;
Submitted (OCT-1997) to the EMBL/GenBank/DDBJ databases.
                                                                       109 AA
                                                                       PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Burkitt's lymphoma receptor-1.";
Nature 391:799-803(1998).
                                                                       STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              NCBI_TaxID=9606;
                                                                       SZ13 HUMAN
                                                                                                              043927
RESULT 1
SZ13_HUMAN
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Chemokine CXCL13/BCA-1.
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                                                                                                             NCBI_TaxID=9544;
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                                                                                                                                                                                                                                                                                                                                                             This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation-the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C., Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M., Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.E., Schnerch A., Schnein J.E., Jones S.J.M., Marra M.A., "Generation and initial analysis of more than 15,000 full-length human
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GO: 0005625; C:soluble fraction; TAS.
GO; GO:0005625; C:soluble fraction; TAS.
GO; GO:000016; F:cell-cell signaling; TAS.
GO; GO:0007267; P:cell-cell signaling; TAS.
GO; GO:0005935; P:chemotaxis; TAS.
GO; GO:000595; P:immune response; TAS.
GO; GO:0006955; P:immune response; TAS.
InterPro; IPR001473; C-X-/Interlan_8.
InterPro; IPR001081; Chemokine_INB.
InterPro; IPR001089; CXC_chmkine_smll.
                                                                                                      and mouse cDNA sequences.";
Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903 (2002).
-!-FUNCTION: Chemocatcive for B lymphocytes but not for T-
lymphocytes, monocytes and but to BLR1/CXCR5.
-!- SUBCELLULAR LOCATION: Secreted.
-!- SIBCELLULAR LOCATION: Secreted.
-!- TISSUE SPECIFICITY: Highest levels in liver, followed by spleen,
lymph node, appendix and stomach. Low levels in salivary gland,
mammary gland and fetal spleen.
-!- SIMILARITY: Belongs to the intercrine alpha (chemokine CXC)
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Last sequence update)
Last annotation update)
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EMBL; AJ002211; CAA05250.1; -.
EMBL; AF029894; AAC17980.1; -.
EMBL; BC012589; AAH12589.1; -.
HSSP; P10145; 3118.
Genew; HGNC:10639; CXCL13.
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01-MAR-2003
01-MAR-2003
01-MAR-2004
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Best Local 9
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Q8HYN9
ID Q8HYI
AC Q8HYI
DT 01-M
DT 01-M
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Gaps
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MEDIJNE=22123042; PubMed=12126650; DOI=10.1006/cyto.2002.0875;
Basu S., Schaefer T.M., Ghosh M., Fuller C.L., Reinhart T.A.;
Basu S., Schaefer T.M., Gequencing of 25 different rhesus macaque chemokine cDNAs reveals evolutionary conservation among C, CC, CXC, AND CX3C families of chemokines.";
Cytokine 18:140-148(2002).

EMBL; AF449284; AAN76087.1; -.
HSSP; P02776; 1F9S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            MEDLINE=22308854; PubMed=12421572; DOI=10.1006/cyto.2002.1973; Pachner A.R., Dail D., Narayan K., Dutta K., Cadavid D.; "Increased expression of B-lymphocyte chemoattractant, but not proinfifammatory cytokines, in muscle tissue in rhesus chronic Lyme
Macaca mulatta (Rhesus macaque).
Eukaryota, Metazoa, Chordata, Craniata, Vertebrata; Euteleostomi;
Mammalia, Eutheria, Primates, Catarrhini, Cercopithecidae;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Cercopithecidae,
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Submitted (APR-2003) to the EMBL/GenBank/DDBJ databases.
EMBL, AV288834; AAP41999.1; --
GO, GO:000576; C:extracellular; IEA.
GO, GO:0008009; F:chemokine activity; IEA.
GO; GO:006855; P:immune response; IEA.
Interpro; IPR01811; Chemokine_ILB.
Fram; PF00048; ILB; I.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PRINTS; PRO0416; INTERLEUKINB.
PRINTS; PRO0437; SWALLCYTKCXC.
SMART; SM00199; SCY; 1.
SEQUENCE 109 AA; 12614 MW; DDB6F9C947DCDECD CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             68 AA; 8030 MW; 46E3D3BB4042EDF2 CRC64;
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01-OCT-2003 (TrEMBLrel. 25, Last sequence update)
01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                        GO; GO:0005576; C:extracellular; IEA.
GO; GO:0008009; F:chmenkine activity; II
GO; GO:0006955; P:immune response; IEA.
InterPro; IPR002473; C-X-C/Interlkn. B.
InterPro; IPR001811; Chemokine ILB.
InterPro; IPR001089; CXC_
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           17.4%; Score 19;
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Macaca mulatta (Rhesus macaque)
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Best Local Similarity
Local 23; Conservative
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Cytokine 19:297-307(2002).
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TISSUE=Human bone marrow;

TISSUE=Human bone marrow;

Wamburte R., Haubner D., Mewes H.W., Weil B., Amid C., Osanger A.,

Fobo G., Han M., Wiemann S.;

L. Submitted (JUN-2003) to the EMBL/GenBank/DDBJ databases.

R EMBL; BX53767; CAD97830.1; -..

R HSSP; p66400; IGUN.

R InterPro; IPR001028; Cyclin.

R InterPro; IPR001028; Cyclin.

R InterPro; IPR0011028; RB A.

R InterPro; IPR00119; RB A.

R Pfam; PF01859; RB A; 1.

R Pfam; PF01857; RB B; 1.

R SWART; SM00385; CYCLIN; 1.

W Hypothetical protein.

O SEQUENCE 849 AA; 95258 MW; DA7274F2B3A9C117 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                 Gaps
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MEDLINE=94074895; PubMed=8253383;
Li Y., Graham C., Lacy S., Duncan A.M.V., Whyte P.;
Li Y., Graham C., Lacy S., Duncan A.M.V., Whyte P.;
The ademovirus ElA-associated 130-kD protein is encoded by a member of the retinoblastoma gene family and physically interacts with
             Gaps
                                                                                                                  01-0CT-2003 (TrEMBLrel. 25, Created)
01-0CT-2003 (TrEMBLrel. 25, Last sequence update)
01-0CT-2003 (TrEMBLrel. 25, Last sequence update)
01-0CT-2003 (TrEMBLrel. 25, Last annotation update)
Hypotherical protein DKFZp781K2028.
Name=DKFZp781K2028;
Name=DKFZp781K2028;
Name=DKFZp781K2028;
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Bukaryota; Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
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01-NOV-1995 (Rel. 39, Last sequence update)
25-OGT-2004 (Rel. 45, Last annotation update)
Retinoblastoma-like protein 2 (130 kDa retinoblastoma-associated
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           Indels
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 Pred. No. 5.4e-12;
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                                                                                                     849 AA
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100.0%; Pred. No. 2.3
tive 0; Mismatches
           Mismatches
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                                           14 PRGNGCPRKEIIVWKKNKS 32
           ;
                               55 PRGNGCPRKEIIVWKKNKS 73
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protein) (PRB2) (P130) (RBR-2)
100.0%;
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           19; Conservative
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  Best Local Similarity
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SEQUENCE FROM N.A.
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RBL2_HUMAN
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           Matches
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | FINAL STATE TYR-672. | MEDINE-2332103; PubMed=1243515; DOI=10.1101/gad.1011202; MEDLINE-2323103; PubMed=1243515; DOI=10.1101/gad.1011202; Tadesco D., Lukas J., Reed S.I.; The pRb-related protein p130 is regulated by phosphorylation-dependent proteolysis via the protein-ubiquitin ligase SCF(Skp2)."; Genes Dev. 16:2946-2957(2002)."; Genes Dev. 16:2957(2002)."; Genes Dev. 16:2957(2002).
                                                 Hannon G.J., Demetrick D., Beach D.; "Isolation of the Rb-related pl30 through its interaction with CDK2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          -!- PTM: Phosphorylation on Ser-672 in G1 leads to its ubiquitin-dependent proteolysis.
-!- MISCELLANEOUS: G0-restricted expression.
-!- SIMILARITY: Belongs to the retinoblastoma protein (RB) family.
                                                                                                                                                                                                                                                                                                        SEQUENCE OF 58-1139 FROM N.A.
MEDLINE=93368960; PubMed=8361765;
MARYOL X., Grana X., Baldi A., Sang N., Hu Q., Giordano A.;
"Cloning of a new member of the retinoblastoma gene family (pRb2)
which binds to the ElA transforming domain.";
Oncogene 8:2561-2566(1993).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TISSUB=Placenta; MEDLINE=96209782; PubMed=8643454; DOI=10.1073/pnas.93.10.4629; MEDLINE=96209782; Paldi A., Boccia V., Claudio P.P., de Luca A., Giordano A.; "Genomic structure of the human retinoblastoma-related Rb2/pl30
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SMART; SM0385; CYCLIN; 2.
SMART; SM0385; CYCLIN; 2.
Phosphorylation: Transcription regulation.
DOWAIN 417, 1024 Pocket; binds ElA.
DOWAIN 417 616 Domain A.
Fig. Spacer.
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InterPro; IPR011028; Cyclin_like.
InterPro; IPR002720; RB_A.
InterPro; IPR002719; RB_B.
Pfam; PF01858; RB_A; 1.
Pfam; PF01857; RB_B; 1.
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Poly-Ala.
Poly-Glu.
Poly-Glu.
MEDLINE=94074896; PubMed=8253384;
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EMBL; S67111; AAB2927.1; -.
EMBL; X74594; CAA52671.1; -.
EMBL; U53220; AAC50479.1; -.
                                                                                                                                                         and cyclins.";
Genes Dev. 7:2378-2391(1993)
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HSSP; P06400; 1GUX.
TRANSFAC; T01608; -.
Genew; HGNC:9894; RBLZ.
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970 RSSSTLPVP 978
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SECG_STRCO
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Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
Altachul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Altschul S.F., Zeeberg B., Bonaldo M.F., Carninci P., Frange C.,
Brownstein M.J., Usdin T.B., Tooshiyuki S., Carninci P., Frange C.,
Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
Allalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
Richards S., Worley W. Sodergren E.J., Lu X., Gibbs R.A.,
Rahey J., Helton B., Ketteman M., Madan A., Young A.C., Shevchenko Y., Boutfard G.G.,
Mhiting M., Madan A., Young A.C., Shevchenko Y., Boutfard G.G.,
Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
Carimwood J., Schmutz J., Myers R.M., Butterfield Y.S.,
Jones S.J., Marra M.A.,
T. "Generation and initial analysis of more than 15,000 full-length human
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Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo
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                                                                                                                                         Length 1139;
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672 672 Phosphoserine.

37 37 S -> P (in Ref. 2).

204 A -> P (in Ref. 1 and 4).

206 206 V -> M (in Ref. 3).

1139 Aa; 128356 MW; ADFBD300E9E1359 CRC64;
                                                                                                                                                                                    0; Indels
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Submitted (JUL-2002) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                01-OCT-2002 (TrEMBLrel. 22, Last sequence update) 01-MAR-2003 (TrEMBLrel. 23, Last annotation update) Retinoblastoma-like 2 (P130).
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                                                                                                                                       8.3%; Score 9; DB 1;
100.0%; Pred. No. 3.1;
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                                                                                                                                       Query Match
Best Local Similarity 100.0%; Pred. No. 3...
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InterPro; IPR011028; Cyclin.
InterPro; IPR002720; RB A.
InterPro; IPR002730; RB B.
Pfam; PP01858; RB A; I.
Pfam; PP01857; RB B; I.
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HSSP; P06400; 1GUX.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Homo sapiens (Human).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               NCBI_TaxID=9606;
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CONFLICT
CONFLICT
                                                                   CONFLICT
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                                                                                                                                                                                                                                                                                                                                                                                         STRAIN=A3(2) / M145,
MEDLINE-2199410; PubMed=12000953; DOI=10.1038/417141a;
MEDLINE-2199410; PubMed=12000953; DOI=10.1038/417141a;
Bencley S. D., Chater K.F., Cerdeno-Tarraga A.-M., Challis G.L.,
Thomson N.R., James K.D., Harris D.E., Quail M.A., Kieser H.,
Harper D., Bateman A., Brown S., Chandra G., Chen C. W., Collins M.,
Cronin A., Fraser A., Goble A., Hidalgo J., Hornsby T., Howarth S.,
Huang C.-H., Kieser T., Larke L., Murphy L.D., Oliver K., O'Neil S.,
Rabbinowited E., Rajandream M.A., Rutherford K.M., Rutter S.,
Seeger K., Saunders D., Sharp S., Squares R., Squares S., Taylor K.,
Warren T., Wietzorrek A., Woodward J.R., Barrell B.G., Parkhill J.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     "Complete genome sequence of the model actinomycete Streptomyces coelicolor A3(2).";
Nature 417:141-147(2002).
-!- FUNCTION: Involved in protein export. Participates in an early event of protein translocation (By similarity).
-!- SUBCELLULAR LOCATION: Integral membrane protein (By similarity).
-!- SIMILARITY: Belongs to the secG family.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Gaps
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SRCGEXPORT.
TIGRRAMs; TIGRO0810; SecG; 1.
Complete protecome; Protein transport; Translocation; Transmembrane.
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                                                                                                                                                                                                                                Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
Streptomycineae; Streptomycetaceae; Streptomyces.
NCBI_TaxID=1902;
                                                                                                                                           Protein export membrane protein secG.
Name=secG, OrderedLocusNames=SCO1944; ORFNames=SCC54.04c;
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100.0%; Pred. No. 3.6;
tive 0; Mismatches 0; Indels
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Last sequence update)
Last annotation update)
                                                  30-MAY-2000 (Rel. 39, Created)
30-MAY-2000 (Rel. 39, Last sequence update)
25-OCT-2004 (Rel. 45, Last annotation update)
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102 AA
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PRT;
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01-MAR-2001 (TrEMBLrel. 16,
01-MAR-2001 (TrEMBLrel. 16,
01-JUN-2003 (TrEMBLrel. 24,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   nterPro; IPR004692; SecG.
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Best Local Similarity 10v...
8; Conservative
STANDARD;
                                                                                                                                                                                                          Streptomyces coelicolor.
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Length 112;

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7.3%; Score 8; DB 2;
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                          8; Conservative
                                                                                                                                               PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Conservative
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                                                   7 SLLLMLLV 14
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            Best Local Similarity
Matches 8; Conserv
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es 8; Conserv
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                                                                                                                                                                                                                                                                                                                                                                             annotation.
                                                                             47
Query Match
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Q8LBD9
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MEDLINE=22608306; PubMed=12692562;
MEDLINE=22608306; PubMed=12692562;
Stead H., Ishikawa J., Hanamoto A., Shinose M., Kikuchi H., Shiba T., Sakaki Y., Hattori M., Omura S.;
"Complete genome sequence and comparative analysis of the industrial microorganism Streptomyces avermitilis.";
Nat. Biotechnol. 21:526-531(2003).
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STRAIN=MA-4680 / ATCC 31267 / NCIMB 12804 / NRRL 8165;
STRAIN=MA-4680 / ATCC 31267 / NCIMB 12804 / NRRL 8165;
MEDLINB=21477403; PubMed=11572948; DOI=10.1073/pnas.211433198;
Omutra S., Ikeda H., Ishikawa J., Hanamoto A., Takahashi C.,
Shinose M., Takahashi Y., Horikawa H., Nakazawa H., Osonoe T.,
Kikuchi H., Shiba T., Sakaki Y., Hattori M.;
"Genome sequence of an industrial microorganism Streptomyces avermitiis: deducing the ability of producing secondary
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
Streptomycineae; Streptomycetaceae; Streptomyces.
NCBI TaxID=33903;
                                     Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
                                                                                                                                                                                                                                                                  Length 102
                                                                                                                                                                                                                                                                                             0; Indels
                                                                                                     Palacin A., Parro V., Mellado R.P.;
Submitted (SEP-1999) to the EMBL/GenBank/DDBJ databases.
EMBL; AF182189; AAG16895.1; -
                                                                                                                                           GO; GO:0016021; C:integral to membrane; IEA.
GO; GO:0015450; F:protein translocase activity; IEA.
GO; GO:0003306; P:protein secretion; IEA.
InterPro; IRR004692; SecG.
Pfam; PF03840; SecG; 1.
PRINTS; PR01651; SECGENDORT.
TIGREPANS; TIGR00810; SecG; 1.
SEQUENCE 102 AA; 10760 MW; 86A498BA0ED00BAA CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                          01-JUN-2003 (TrEMBLrel. 24, Created)
01-JUN-2003 (TrEMBLrel. 24, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
Putative SecG-like integral membrane protein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Proc. Natl. Acad. Sci. U.S.A. 98:12215-12220(2001)
                                           Streptomycineae; Streptomycetaceae; Streptomyces.
                                                                                                                                                                                                                                                                 7.3%; Score 8; DB 2;
100.0%; Pred. No. 3.6;
tive 0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                    112 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Name=secG; OrderedLocusNames=SAV6299; Streptomyces avermitilis.
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TIGREAMS; TIGR00810; SecG; 1.
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Pfam; PF03840; SecG; 1.
                                                                                                                                                                                                                                                    Query Match
Best Local Similarity 10v..
                                                                                                                                                                                                                                                                                                                                                                                                                   PRELIMINARY;
                         Streptomyces lividans.
                                                                                                                                                                                                                                                                                                                                        7 SLLLMLLV 14
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                                                                                          SEQUENCE FROM N.A.
            Name=secG;
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Q829W0
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01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
01-OCT-2002 (TrEMBLrel. 26, Last sequence update)
01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
Putative proline-rich protein.
Arabidopsis thaliana (Mouse-ear cress).
Arabidopsis thaliana (Mouse-ear cress).
Sukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnollophyta; eudicotyledons; core eudicots; rosids;
eurosids II; Brassicales; Brassicaceae; Arabidopsis.
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Bukaryota, Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; rosids;
eurosids II; Brassicales; Brassicaceee; Arabidopsis.
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Yamada K., Chan M.M., Chang C.H., Dale J.M., Heuan V.W., Lee J.M.,
Onodera C.S., Quach H.L., Tang C.C., Toriumi M., Wong C., Wu H.C.,
Yu G., Yuan S., Chen H., Cheuk R., Jones T., Kim C.J., Nguyen M.,
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MEDILINE=22088475; PubMed=12093376;
Haas B.J., Volfovsky N., Town C.D., Troukhan M., Alexandrov
Feldmann K.A., Flavell R.B., White O., Salzberg S.L.;
"Full-length messenger RNA sequences greatly improve genome
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Brover V., Troukhan M., Alexandrov N., Lu Y.-P., Flavell R., Feldmann K.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 239;
                         0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Submitted (MAR-2002) to the EMBL/GenBank/DDBJ databases.
EMBL; AY087270; AAM64824.1; -.
GO; GO:005198; F:structural molecule activity; IEA.
InterPro; IPR000535; MSP.
InterPro; IPR00862; PapD-like.
InterPro; IPR008162; Pantne_S.
Pfam; PF00635; Morile_Sperm; 1.
PROSITE; PS00012; PHOSPHOPANTETHEINE; UNKNOWN_1.
SEQUENCE 239 AA; 26327 MW; 3D57A5E8B0EEC76B CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                01-JUN-2003 (TrEMBLrel. 24, Created)
01-JUN-2003 (TrEMBLrel. 24, Last sequence update)
01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
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100.0%; Pred. No. o...
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100.0%; Pred. No. 4; ive 0; Mismatches
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PRELIMINARY;
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                                                                                                                                                      Escherichia coli 06.
                                                                                                                                                                                                                                                       SEQUENCE FROM N.A.
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  Q8FK19
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SEQUENCE FROM N.A.

STRAIN=PCA / ATCC 51573;

STRAIN=PCA / ATCC 51573;

STRAIN=PCA / ATCC 51573;

STRAIN=PCA / ATCC 51573;

Methe B.A., Nelson K.B., Eisen J.A., Paulsen I.T., Nelson W.C.,

Heidelberg J.F., Wu D., Wu M., Ward N.L., Beanan M.J., Dodson R.J.,

Madugu R., Brinkac L.M., Daugherty S.C., DeBoy R.T., Durkin A.S.,

Gwinn M.L., Kolonay J.F., Sullivan S.A., Haft D.H., Selengut J.,

Davidsen T.M., Zafar N., White O., Tran B., Romero C., Forberger H.A.,

Weidman J.F., Khouri H.M., Feldblyum T.V., Utterback T.R.,

Wan Aken S.B., Lovley D.R., Fraser C.M.,

"Genome of Geobacter sulfurreducens: metal reduction in subsurface
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
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Geobacteraceae; Geobacter.
Palm C.J., Shinn P., Southwick A., Tripp M.G., Wu T., Davis R.W.,
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GO:0015087; F:cobalt ion transporter activity; IEA.
GO:0015095; F:magnesium ion transporter activity; IEA.
GO:0006824; P:cobalt ion transport; IEA.
GO:0015693; P:magnesium ion transport; IEA.
                                                                                                                                                                                                                                                     Length 239;
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              Ecker J.R., Theologis A.;
Submitted (NOV-2002) to the EMBL/GenBank/DDBJ databases.
EMBL; BT001917; AAN11916.1; -.
                                                                       GO, GO:0005198; F:structural molecule activity; IEA.
InterPro; IPR000535; MSP.
InterPro; IPR006162; Papholike.
InterPro; IPR006162; Papholike.
InterPro; IPR006162; Papholike.
PROSITE; PS00012; MSP; 1.
PROSITE; PS00012; PHOSPHORANTETHEINE; UNKNOWN 1.
SEQUENCE 239 AA; 26374 MW; 2CA7AB2B7DF1E54B CRC64;
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05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
Magnesium and cobalt transport protein CorA.
Name-corA-1; OrderedLocusNames=GSU1399;
                                                                                                                                                                                                                                                     Query Match 7.3%; Score 8; DB 2; Best Local Similarity 100.0%; Pred. No. 8.1; Matches 8; Conservative 0; Mismatches
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100.0%; Pred. No. 12;
iive 0; Mismatches
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InterPro; IPR002523; Mg2+_transptCorA.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          environments.";
Science 302.1967-1969(2003).
EMBL; AE011180; AAR34775.1; -
TIGR; GSU1399; -
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Matches 8; Conservative
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RESULT 13

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            STRAIN=06.HI / CFT073 / ATCC 700928;
MEDLINE=22388234; PubMed=12471157; DOI=10.1073/pnas.252529799;
Welch R.A., Burland V., Plunkett G. III, Redford P., Rossch P.,
Rasko D., Buckles E.L., Liou S.-R., Boutin A., Hackett J., Stroud D.,
Mayhew G.F., Rose D.J., Schwartz D.C., Perna N.T.,
Mobley H.L.T., Donnenberg M.S., Blattner F.R.;
"Extensive mosaic structure revealed by the complete genome sequence
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01-007-2002 (TrEMBLrel. 22, Last sequence update)
01-007-2002 (TrEMBLrel. 28, Last sequence update)
CG3576-PA (CG3576-pb) (LD18904p) (Longevity protein).
Name-Lagl; Synonyms-CG15898, lon; ORFNames=CG3576;
Drosophila melanogaster (Fruit fly).
Eukaryota, Metazoa, Arthropoda; Hexapoda; Insecta; Pterygota;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
Ephydroidea; Drosophilidae; Drosophila.
NCBI_TaxID=7227;
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                                                                                                                         u1-mak-2003 (TrEMBLrel. 23, Last sequence update) 01-0CT-2003 (TrEMBLrel. 25, Last annotation update) Isochorismate synthase entC (EC 5.4.99.6).
Mame-entC; OrderedLocusNames=c0680;
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395 AA
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100.0%; Pred. No. 13;
ive 0; Mismatches
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01-MAR-2003 (TrEMBLrel. 23, Last seq
01-OCT-2003 (TrEMBLrel. 25, Last ann
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NCBI_TaxID=217992;
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Best Local Similarity 100.0
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RA Cherry J.W., Cawley S., Dahlke C., Davenport L.B., Davies P.,

A dodgon K., Delcher A., Deng Z., Mays A.D., Dew I., Dietz S.M.,

RA Dodgon K., Doup L.B., Downes M., Dugan-Rocha S., Dunkov B.C., Dunn P.,

RA Durbin K.J., Evangelista C.C., Ferraz C., Ferriera S., Fleischmann W.,

RA Fosler C., Gabrielian A.E., Garg N.S., Gelbart W.M., Glasser K.,

RA Harris N.L., Harvey D., Heinan T.J., Wei M.H., Ibegwam C.,

RA Harris N.L., Harvey D., Heinan T.J., Wei M.H., Ibegwam C.,

RA Jalai M., Kalush F., Karpen G.H., Ke Z., Kannison J.A.,

RA Lasko P., Lei Y., Levitsky A.A., Li J., Li Z., Liang Y., Lin X.,

RA Lasko P., Lei Y., Levitsky A.A., Li J., Li Z., Liang Y., Lin X.,

RA Martei B. W. Morbosh T.C., Morris J., Moshrefi A.,

RA Mount S.M., Moy M., Murphy B., Murphy L., Muzny D.M., Nelson D.L.,

RA Palazolo M., Pittman G.S., Pan S., Pollard J., Puri, Pacle D.M.,

RA Reinert K., Remington K., Saunders R.D., Scheeler F., Shen H.,

Rhe B.C., Siden-Kiamos I., Simpson M., Skupski M.P., Smith T.,

Rhu B.C., Siden-Kiamos I., Simpson M., Strong R., Sun E.,

Syriskas R., Tector C., Turner R., Venter E., Wang A.H., Wang X.,

Williams S.M., Woodager, Worley K.C., Wu D., Yang S., Yao Q.A., Ye J.,

RA Zheng X.H., Zhong F.N., Zhan M., Zhou X., Zhu S., Zhu X., Smith H.O.,

RA Zheng X.H., Zhong F.N., Rubin G.M., Venter J.C.;

RA Schence 287:2185-2195(2000).
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"The transposable elements of the Drosophila melanogaster euchromatin:
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Misra S., Crosby M.A., Mungall C.J., Matthews B.B., Campbell K.S.,
Hradecky P., Huang Y., Kaminker J.S., Millburn G.H., Prochnik S.E.,
Smith C.D., Tupy J.L., Whitfied E.J., Bayraktaroglu L., Berman B.P.
Bettencourt B.R., Celniker S.E., de Grey A.D., Drysdale R.A.,
Harris N.L., Richter J., Russo S., Schroeder A.J., Shu S.Q.,
Stapleton M., Yamada C., Ashburner M., Gelbart W.M., Rubin G.M.,
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"Annotation of the Drosophila melanogaster euchromatic genome: a
systematic review.";
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Stapleton M., Brokstein P., Hong L., Agbayani A., Carlson J.,
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Genome Biol. 3:RESEARCH0084-RESEARCH0084(2002).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   melanogaster euchromatic genome sequence.";
Genome Biol. 3:RESEARCH0079-RESEARCH0079(2002).
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Genome Biol. 3:RESEARCH0083-RESEARCH0083(2002)
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PubMed=14960714; DOI=10.1093/nar/gkh258;
Rasko D.A., Ravel J., Oekstad O.A., Helgason E., Cer R.Z., Jiang L., Shores R.A., Fouts D.E., Tourasse N.J., Angiuoli S.V., Kolonay J.F., Nelson W.C., Kolstoe A.-B., Fraser C.M., Read T.D.; T.V., Kolstoe A.-B., Fraser C.M., Read T.D.; The genome sequence of Bacillus cereus ATCC 10987 reveals metabolic adaptations and a large plasmid related to Bacillus anthracis pXO1."; Nucleic Acids Res. 32:977-988 (2004).

FMEL; AE017267; AAS40032.1; TIGR; BCEI101; TIGR;
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                                                                                                                                                                                                                                                                                                                                                                                                         Publicator, 2012. Pagno40918; Lagl.

GO; GO:0016021; C:integral to membrane; IEA.

GO; GO:0005634; C:integral to membrane; IEA.

GO; GO:0003700; F:transcription factor activity; IEA.

GO; GO:0006355; P:regulation of transcription, DNA-dependent; IEA.

InterPro; IPR001356; Homeodox.

InterPro; IPR009057; Homeodomain_like.
Champe M., Chavez C., Dorsett V., Farfan D., Frise E., George R., Goralez M., Guarin H., Li P., Liao G., Miranda A., Mungall C.J., Nunco J., Pacleb J., Paragas V., Park S., Phouanenavong S., Wan K. Yu C., Lewis S.E., Rubin G.M., Celniker S.; Submitted (OCT-2001) to the EMBL/GenBank/DDBJ databases.
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                                                                                                                                                                                         Bauer R., Eckardt F., Lehmann C., Hoch M.;
Submitted (DEC-2001) to the EMBL/GenBank/DDBJ databases.
-!- SUBCELLULAR LOCATION: Nuclear (By similarity).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Bacillus cereus (strain ATCC 10987).
Bacteria; Firmicutes; Bacillales; Bacillaceae; Bacillus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               PROSITE; PS50071; HOMBCBOX_2; 1.
PROSITE; PS50922; TLC; 1.
DNA-binding; Homeobox; Nuclear protein.
SEQUENCE 400 AA; 46350 MW; AB586DB2B8111E62 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     46284 MW; E7664EF1BF3C654F CRC64;
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05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
Sensor histidine kinase, putative (EC 2.7.3.-).
OrderedLocusNames=BCE1101;
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100.0%; Pred. No. 13;
tive 0; Mismatches
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                                                                                                                                                                                                                                                                       EMBL; AE000437; AAF46137.2; ---
EMBL; AY061255; AAL28803.1; --
EMBL; AX066020; AAL57756.1; --
EMBL; AY066020; ALE8.
Intact; Q9W423; --
FlyBase; FBgn0040918; Lag1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          InterPro; IPR006634; TLC.
Pfam; PF00798; LAG1; 1.
PRINTS; PR00024; HOMEOBOX.
SMART; SM00389; HOX; 1.
SMART; SM00724; TLC; 1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             309 SLLLMLLV 316
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Gaps ; 0 0; Indels Best Local Similarity 100.0%; Pred. No. 13; Matches 8; Conservative 0; Mismatches **상** 원

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41 VFIPRRFI 48 ||||||||| 18 VFIPRRFI 25

Search completed: September 21, 2005, 13:44:27 Job time : 178 secs

Sequence 1143, Sequence 3

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Sequence 1720, F Sequence 1795, P Sequence 2059, A Sequence 225, A Sequence 2256, A Sequence 2266, A

Sequence Sequence Sequence Sequence Sequence

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Sequence Sequence

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APPLICANT: LI, HAODONG
APPLICANT: LI, HAODONG
TITLE OF INVENTION: THERAPEUTIC COMPOSITIONS AND METHODS FOR
TITLE OF INVENTION: TREATING DISEASE STATES WITH LEUKOCYTE ADHESION
INHIBITOR-1 (LAI-1), AND CHEMOKINE BETA-11 (CRBETA-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  NUMBER OF SEQUENCES: 16
CORRESPONDENCE ADDRESS:
ADDRESSE: STERNE, KESSLER, GOLDSTEIN & FOX, P.L.L.C.
STREET: 1100 NEW YORK AVENUE, SUITE 600
CITY: WASHINGTON
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      COUNTRY: USA

COUNTRY: USA

ZIP: 20005

COUNTRY: USA

ZIP: 20005

COMPUTER READABLE FORM:

MEDIUM TYPE: Floppy disk

COMPUTER: IBM PC compatible

COMPUTER: PatentIn Release #1.0, Version #1.30

CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/10/247,287

FILING DATE: 20-Sep-2002

PRIOR APPLICATION NUMBER: US/09/635,899

FILING DATE: 11-Aug-2000

APPLICATION NUMBER: US 08/943,336

FILING DATE: 03-OCT-1997

APPLICATION NUMBER: US 06/027,769

FILING DATE: 04-OCT-1996
                                                                                                                                                                                                                                                                  US-10-040-862-1143
US-10-040-862-1151
US-10-040-862-1151
US-10-040-862-1185
US-10-040-862-1200
US-10-040-862-1205
US-10-040-862-1254
US-10-040-862-1355
US-10-040-862-1612
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US-10-040-862-1663
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US-09-796-692-1707
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US-09-796-692-1707
US-09-796-692-2256
US-09-796-692-2256
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ; Sequence 4, Application US/10247287; Publication No. US20030082770A1; GENERAL INFORMATION:
    US-10-247-287-4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      RESULT 1
    Sequence 4, Appli
Sequence 240, App
Sequence 240, App
Sequence 158, Appl
Sequence 2, Appli
Sequence 146, App
Sequence 1494, Ap
Sequence 10, Appl
Sequence 5091, Appl
Sequence 5091, Appl
Sequence 5091, Appl
                                                                                                   September 21, 2005, 13:41:21 ; Search time 63 Seconds
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704.196 Million cell updates/sec
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                                                                                                                                                                                           MKFISTSLLLMLLVSSLSPV......VLRKRSSSTLPVPVFKRKIP 109
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1: /cgn2_6/ptodata/1/pubpaa/US07_PUBCOMB.pep:*

2: /cgn2_6/ptodata/1/pubpaa/PCT_MEW_PUB_pep:*

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17: /cgn2_6/ptodata/1/pubpaa/US10B_PUBCOMB.pep:*

18: /cgn2_6/ptodata/1/pubpaa/US10B_PUBCOMB.pep:*

19: /cgn2_6/ptodata/1/pubpaa/US10B_PUBCOMB.pep:*

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11: /cgn2_6/ptodata/1/pubpaa/US11A_REW_PUB.pep:*

12: /cgn2_6/ptodata/1/pubpaa/US11A_REW_PUB.pep:*

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16: /cgn2_6/ptodata/1/pubpaa/US11A_REW_PUB.pep:*

17: /cgn2_6/ptodata/1/pubpaa/US11A_REW_PUB.pep:*

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                 GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd.
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1 US-10-050-902-240
1 US-10-050-902-240
1 US-10-247-671-158
1 US-10-247-671-158
2 US-10-705-314-2
2 US-10-755-889-146
2 US-10-755-889-146
3 US-10-755-889-146
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Gapop 60.0 , Gapext 60.0
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and is derived by analysis of
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seq length: 200000000
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Maximum DB
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Perfect :
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Result No.

1109876511

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1 MKFISTSLLLMLLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC
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                                                                                                                                                                             61 PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
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                                                                                                                                                61 PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
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Indels
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APPLICANT: Shiffman, Dov
APPLICANT: Porter, Gordon, J.
APPLICANT: Reser, Matthew R.
TITLE OF INVENTION: GENES EXPRESSED IN TREATED FOAM CELLS
FILE REPERBNE: PA-0050 US
CURRENT APPLICATION NUMBER: US/10/247,671
CURRENT FILING DATE: 2002-09-18
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; Pred. No. 1.5e-96;
0; Mismatches 0;
0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 APPLICANT: Frey, Peter
TITLE OF INVENTION: Molecular Antigen Array
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             CURRENT FILING DATE: 2002-01-18
PRIOR PILING DATE: 2001-01-9
PRIOR PILING DATE: 2001-01-9
PRIOR APPLICATION NUMBER: US 60/288,549
PRIOR APPLICATION NUMBER: US 60/326,998
PRIOR FILING DATE: 2001-10-05
PRIOR FILING DATE: 2001-10-05
PRIOR PILING DATE: 2001-10-05
NUMBER OF SEQ ID NOS: 350
SOFTWARE: PATENTIN OFF: 2011-107
SSOFTWARE: PATENTIN OFF: 2011-11-07
SSOFTWARE: PATENTIN OFF: 2011-11-07
SSOFTWARE: PATENTIN OFF: 2011-11-07
SSOFTWARE: PATENTIN OFF: 2.1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    CURRENT APPLICATION NUMBER: US/10/050,898 CURRENT FILING DATE: 2002-01-18
                                                                                                                                                                                                                                                                                                                      Sequence 240, Application US/10050898
Publication No. US20030175711A1
GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Sequence 158, Application US/10247671 Publication No. US20030194721A1 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Luond, Rainer
Staufenbiel, Matthias
                                                                                                                                                                                                                                                                                                                                                                                         APPLICANT: Renner, Wolfgang A. APPLICANT: Bachmann, Martin APPLICANT: Tissot, Alain APPLICANT: Cechner, Patrick APPLICANT: Lechner, Franziska APPLICANT: Sebbel, peter APPLICANT: Ortmann, Rainer
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity 100.0%;
Matches 109; Conservative 0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    FILE REFERENCE: 1700.0190005
Matches 109; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   CRGANISM: Homo sapiens US-10-050-898-240
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APPLICANT:
APPLICANT:
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Pred. No. 1.5e-96;
                                                                FILING DATE: 05-JUN-1996
APPLICATION NUMBER: US 08/464,401
FILING DATE: 05-JUN-1995
APPLICATION NUMBER: US 08/460,987
FILING DATE: 05-JUN-1995
APPLICATION NUMBER: PCT/US95/01780
FILING DATE: 08-FEB-1995
ATONEY/AGENT INFORMATION:
NAME: STEPFE, ERIC K.
REGISTRATION NUMBER: 36,688
REFERENCE/DOCKET NUMBER: 1488.0380006
TELECRANIONICATION INFORMATION:
TELECRANICATION INFORMATION:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        GENERAL INFORMATION:
APPLICANT: Bachmann, Martin
APPLICANT: Tissot, Alain
APPLICANT: Tissot, Alain
APPLICANT: Tissot, Alain
APPLICANT: Dechner, Patrick
APPLICANT: Dechner, Patrick
APPLICANT: Dechner, Pranziska
APPLICANT: Sebbel, Peter
APPLICANT: Posssek, Christine
ITILE OP INVENTION: Molecular Antigen Array
FILE REFERENCE: 1700.0190004
CURRENT FILING DATE: 2002-01-18
PRIOR APPLICATION NUMBER: US 60/262,379
PRIOR APPLICATION NUMBER: US 60/262,379
PRIOR PILING DATE: 2001-01-19
PRIOR APPLICATION NUMBER: US 60/266,998
PRIOR FILING DATE: 2001-00-05
PRIOR FILING DATE: 2001-10-05
PRIOR FILING DATE: 2001-11-07
NUMBER OF SEQ ID NOS: 350
SOFTHARE: Patentin Ver: 2.1
                       FILING DATE: 04-OCT-1996
APPLICATION NUMBER: PCT/US96/09572
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Mismatches
US 08/724,871
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          TOPOLOGY: linear
MOLECULE TYPE: protein
SEQUENCE DESCRIPTION: SEQ ID NO: 4:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Application US/10050902
o. US20030175290A1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           LENGTH: 109 amino acids TYPE: amino acid
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100.0%;
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SEQUENCE CHARACTERISTICS
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Best Local Similarity 100.
Matches 109; Conservative
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                                                                                                                                                                                               Length 109;
                                                                                                                       FEATURE:

NAME/KGY: misc feature

OTHER INFORMATION: Incyte ID No. US20030194721A1 020293CD1
US-10-247-671-158
                                                                                                                                                                                                                              0; Indels
                                                                                                                                                                                               Query Match 100.0%; Score 109; DB 14; Best Local Similarity 100.0%; Pred. No. 1.5e-96; Matches 109; Conservative 0; Mismatches 0;
PRIOR APPLICATION NUMBER: 60/323,784
PRIOR FILING DATE: 2001-09-19
NUMBER OF SEQ ID NOS: 186
SOFTWARE: PERL Program
SEQ ID NO 158
LENGTH: 109
                                                                                                                                                                                                                                                                                                                                                                                                                            ; Sequence 88, Application US/10058270A; Publication No. US20040029114A1
                                                                         LENGTH: 109
TYPE: PRT
ORGANISM: Homo sapiens
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ORGANISM: Homo sapiens
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APPLICANT: Bristol-Myers Squibb Company
TITLE OF INVENTION: POLYNUCLEOTIDES AND POLYPEPTIDES ASSOCIATED WITH THE NF-KB
TITLE OF INVENTION: PATHWAY
FILLE OF INVENTION: PATHWAY
FILLE OF INVENTION: PATHWAY
CURRENT APPLICATION NUMBER: US/10/755,889
CURRENT APPLICATION NUMBER: US. 60/440,068
PRIOR APPLICATION NUMBER: U.S. 60/469,757
PRIOR FILLING DATE: 2003-01-14
PRIOR FILLING DATE: 2003-05-12
NUMBER OF SEQ ID NOS: 823
SOFTWARE: PATENTIN VERSION 3.2
SEQ ID NO 146
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VESULI 6

US-10-700-314-2

Sequence 2, Application US/10700314

Publication No. US20040086975A1

GENERAL INFORMATION:

APPLICANT: GUEGLER, Karl J.

APPLICANT: WILDE, Craig G.

APPLICANT: BELILAMER, Jeffrey J.

TITLE OF INVENTION: CHEMOKINE EXPRESSED IN INFLAMED ADENOID FILE REFERENCE: PF-0025-4 DIV

CURRENT PELLING DATE: 2003-10-28

PRIOR APPLICATION NUMBER: US 09/588,044

PRIOR APPLICATION NUMBER: US 09/588,044

PRIOR APPLICATION NUMBER: US 09/588,044

PRIOR APPLICATION NUMBER: US 09/203,235

PRIOR FILING DATE: 1908-12-01

PRIOR FILING DATE: 1994-12-07

PRIOR FILING DATE: 1994-12-07

PRIOR FILING DATE: 1994-12-07

NUMBER OF SEQ ID NOS: 2

SOFTWARE: PERL PROGRAM

SEQ ID NO 2

LENGTH DATE: PROGRAM
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ; NAME/KEY: misc_feature
; OTHER INFORMATION: Incyte ID No: 020293CD1
US-10-700-314-2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ; Sequence 146, Application US/10755889; Publication No. US20040171823A1; GENERAL INFORMATION:
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US-10-755-889-146
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RESULT 10
US-10-756-149-5091
Sequence 5091, Application US/10756149
Sequence 5091, Application US/10756149
Sequence 5091, Application No. US20050181375A1
GENERAL INFORMATION:
APPLICANT: Albert
SPELICANT: Albert
TITLE OF INVENTION: NOVEL METHODS OF DIAGNOSIS OF METASTATIC CANCER
TITLE OF INVENTION: METHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER
TITLE OF INVENTION: METHODS OF SCREENING FOR MODULATORS OF METASTATIC CANCER
CURRENT APPLICATION NUMBER: US/10/756,149
CURRENT FILING DATE: 2004-01-12
NUMBER OF SEQ ID NOS: 5818
SOFTWARES: PatentIn version 3.2
SEQ ID NO 5091
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                                                                                             61 PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
                                                                                                                                61 PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLFVEVFKRKIP 109
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Bublication No. US20020198362A1

GENERAL INFORMATION:
APPLICANT: Gaiger, Alexander
APPLICANT: Gaiger, Paul A.
APPLICANT: Gaiger, Paul A.
APPLICANT: Algate, Paul A.
TITLE OF INVENTION: COMPOSITIONS AND METHODS FOR THE TITLE OF INVENTION: HEMATOLOGICAL MALIGNANCIES
TITLE OF INVENTION: COMPOSITIONS AND METHODS FOR THE TITLE OF INVENTION: HEMATOLOGICAL MALIGNANCIES
CURRENT PELLOATION NUMBER: US/09/796,692
CURRENT FILING DATE: 2000-03-01
PRIOR APPLICATION NUMBER: 60/186,126
PRIOR PELLOATION NUMBER: 60/190,479
PRIOR FILING DATE: 2000-04-27
PRIOR APPLICATION NUMBER: 60/200,303
PRIOR PELLOATION NUMBER: 60/200,779
PRIOR PELLOATION NUMBER: 60/200,779
PRIOR PELLOATION NUMBER: 60/200,779
PRIOR APPLICATION NUMBER: 60/200,779
PRIOR PELLOATION NUMBER: 60/200,779
PRIOR APPLICATION NUMBER: 60/200,999
PRIOR PELLICATION NUMBER: 60/200,999
PRIOR PELLICATION NUMBER: 60/200,999
PRIOR PELLICATION NUMBER: 60/200,999
PRIOR PELLICATION NUMBER: 60/200,999
PRIOR PELLING DATE: 2000-05-04
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100.0%; Pred. No. 1.5e-96;
iive 0; Mismatches 0;
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Matches 109; Conservative
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US-10-756-149-5091
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| Sequence 10, Application US/10510121
| Publication No. US20050119174A1
| GENERAL INFORMATION:
| APPLICANT: ICHIHARA, Junji
| FILE REFERENCE: 231060
| FILE REFERENCE: 231060
| FILE REFERENCE: 2004-10-04
| PRIOR APPLICATION NUMBER: US/10/510,121
| PRIOR APPLICATION NUMBER: PCT/JP03/04260
| PRIOR APPLICATION NUMBER: PCT/JP03/04260
| PRIOR FILING DATE: 2003-04-03
| NUMBER OF SEQ ID NOS: 10
| SEQ ID NOS: 10
| SEQ ID NO 10
| SEQ ID NO 10
                                                                                                                                                                                                                                                                                                                                                                                                    APPLICANT: Aliz, Natasha
APPLICANT: Ginsburg, Wendy M.
APPLICANT: Ginsburg, Wendy M.
APPLICANT: Ginsburg, Wendy M.
APPLICANT: Zlotnik, Albert
TITLE OF INVENTION: Methods of Diagnosis of Soft Tissue Sarcoma Modulators
TITLE OF INVENTION: Methods for Screening for Soft Tissue Sarcoma Modulators
FILE REFERENCE: 05882.0193.NPUS01
CURRENT APPLICATION NUMBER: US/10/723,860
CURRENT APPLICATION NUMBER: 06/429,739
PRIOR APPLICATION NUMBER: 60/429,739
PRIOR APPLICATION NUMBER: 60/429,739
NUMBER OF SEQ ID NOS: 8393
SOFTWARE: Patentin version 3.2
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   1 MKFISTSLLLMLLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC
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                                                                                                                                   61 PRKEIIVWKKNKSIVCVDPQAEWIQRAMEVLRKRSSSTLPVPVFKRKIP 109
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                                                                                                                                                                                                                                                                                                       Sequence 1494, Application US/10723860
Publication No. US20040253606A1
GENERAL INFORMATION:
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ORGANISM: Homo sapiens
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ORGANISM: Homo sapiens
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Best Local Similarity
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GREACH LIVEWANTION:

APPLICANT: Algate, Paul A.

APPLICANT: Mannion, Jane

TITLE OF INVENTION: COMPOSITIONS AND METHODS FOR THE DETECTION, DIAGNOSIS AND THERAPY

TITLE OF INVENTION: UNMER: 105/09/196,692

CURRENT APPLICATION NUMBER: 60/186,126

PRIOR APPLICATION NUMBER: 60/180,479

PRIOR PLING DATE: 2000-03-17

PRIOR PLING DATE: 2000-04-28

PRIOR PLING DATE: 2000-05-01

PRIOR PLING DATE: 2000-05-01

PRIOR PLING DATE: 2000-05-04

PRIOR PLING DATE: 2000-06-04

PRIOR PLING DATE: 2000-08-04

PRIOR PLING DATE: 2000-08-04

PRIOR PLING DATE: 2000-08-04

PRIOR PLING DATE: 2000-08-03

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                   0; Indels
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100.0%; Pred. No. 1.6e-96;
iive 0; Mismatches 0;
                   0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ; Sequence 1167, Application US/09796692; Publication No. US20020198362A1; GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ; Sequence 1185, Application US/09796692; Publication No. US20020198362A1
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Best Local Similarity 100.0
Matches 109; Conservative
                   Matches 109; Conservative
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ORGANISM: Homo sapiens
US-09-796-692-1167
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APPLICANT: Galger, Alexander
APPLICANT: Galger, Alexander
APPLICANT: Galger, Alexander
APPLICANT: Galger, Alexander
APPLICANT: Algate, Paul A
TITLE OF INVENTION: COMPOSITIONS AND METHODS FOR THE DETECTION, DIAGNOSIS AND THERAPY
TITLE OF INVENTION: HEMATOLOGICAL MALIGNANCIES
TITLE OF INVENTION: HEMATOLOGICAL GALGE
CURRENT APPLICATION NUMBER: 60/186,126
PRIOR APPLICATION NUMBER: 60/186,126
PRIOR APPLICATION NUMBER: 60/180,479
PRIOR APPLICATION NUMBER: 60/200,303
PRIOR PILING DATE: 2000-03-01
PRIOR PELING DATE: 2000-03-07
PRIOR PELING DATE: 2000-03-07
PRIOR PELING DATE: 2000-03-07
PRIOR PELING DATE: 2000-05-07
PRIOR PILING DATE: 2000-05-07
PRIOR PELING DATE: 2000-05-07
PRIOR PELING DATE: 2000-05-07
PRIOR PELING DATE: 2000-05-07
PRIOR PELING DATE: 2000-07-04
PRIOR PELING DATE: 2000-07-04
PRIOR PELING DATE: 2000-07-04
PRIOR PELING DATE: 2000-08-07
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Pred. No. 1.6e-96;
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100.0%; Pred. No. 1.6e-96;
ive 0; Mismatches 0;
; PRIOR FILING DATE: 2000-08-03

; PRIOR APPLICATION NUMBER: 60/223,416

; PRIOR FILING DATE: 2000-08-04

; PRIOR FILING DATE: 2000-08-07

; NUMBER OF SEQ ID NOS: 9597

; SOFTWARE: FastSEQ for Windows Version 3.0

; SEQ ID NO 1143
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Publication No. US20020198362A1
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Matches 109; Conservative
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; ORGANISM: Homo sapiens
US-09-796-692-1143
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; ORGANISM: Homo sapiens
US-09-796-692-1151
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Best Local Similarity
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; Pred. No. 1.6e-96;
0; Mismatches 0;
PRIOR APPLICATION NUMBER: 60/200,303
PRIOR FILING DATE: 2000-04-28
PRIOR FILING DATE: 2000-04-28
PRIOR FILING DATE: 2000-04-28
PRIOR FILING DATE: 2000-05-01
PRIOR FILING DATE: 2000-05-01
PRIOR PILING DATE: 2000-05-01
PRIOR PILING DATE: 2000-05-04
PRIOR PILING DATE: 2000-05-22
PRIOR PILING DATE: 2000-05-22
PRIOR FILING DATE: 2000-05-22
PRIOR PILING DATE: 2000-05-22
PRIOR PILING DATE: 2000-06-31
PRIOR PILING DATE: 2000-08-03
PRIOR PILING DATE: 2000-08-03
PRIOR PILING DATE: 2000-08-03
PRIOR PILING DATE: 2000-08-04
PRIOR PILING DATE: 2000-08-07
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100.0%;
Best Local Similarity 100.0%;
Matches 109; Conservative 0;
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US-09-796-692-1200
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ORGANISM:
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                                    APPLICANT: Algater, Paul A.
APPLICANT: Algater, Paul A.
APPLICANT: Algater, Paul A.
APPLICANT: Algater, Paul A.
APPLICANT: Algater, Paul A.
APPLICANT: Algater, Paul A.
APPLICANT: Mannion, Jane
TITLE OF INVENTION: COMPOSITIONS AND METHODS FOR THE DETECTION, DIAGNOSIS AND THERAPY
FILLE OF INVENTION: UNMERS: US/O9/796,592
CURRENT APPLICATION NUMBER: 60/186,126
REIGHT FILLING DATE: 2000-03-1
REIGHT FILLING DATE: 2000-03-1
REIGHT FILLING DATE: 2000-04-27
REIGHT APPLICATION NUMBER: 60/200,545
REIGHT APPLICATION NUMBER: 60/200,545
REIGHT APPLICATION NUMBER: 60/200,999
REIGHT APPLICATION NUMBER: 60/202,416
REIGHT APPLICATION NUMBER: 60/202,416
REIGHT APPLICATION NUMBER: 60/223,416
REIGHT APPLICATION NUMBER: 60/223,416
REIGHT APPLICATION NUMBER: 60/223,416
REIGHT REILING DATE: 2000-09-03
REIGHT REILING DATE: 2000-09-04
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Publication No. US20020199362A1

GENERAL INFORMATION:
APPLICANT:
APPLICANTION:
CURRENT APPLICATION NUMBER: US/09/796,692
CURRENT APPLICATION NUMBER: 00/166,126
PRIOR FILING DATE: 2000-03-01
PRIOR APPLICATION NUMBER: 60/190,479
PRIOR FILING DATE: 2000-03-17
PRIOR APPLICATION NUMBER: 60/200,545
PRIOR FILING DATE: 2000-04-27
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100.0%; Score 109; DB 9;
Best Local Similarity 100.0%; Pred. No. 1.6e-96;
Matches 109; Conservative 0; Mismatches 0;
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SOFTWARE: FastSEQ for Windows Version 3.0
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US-09-796-692-1185
                  GENERAL INFORMATION:
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US-09-796-692-1200
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Query Match 100.0%; Score 109; DB 1; Length 109; Best Local Similarity 100.0%; Pred. No. 6e-98; Matches 109; Conservative 0; Mismatches 0; Indels
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; CLONE: 20293
US-08-352-324A-2
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189.227 Million cell updates/sec
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Sequence 4,
Sequence 4,
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1 MKFISTSLLLMLLVSSLSPV.....VLRKRSSSTLPVPVFKRKIP 109
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Sequence 9,
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Sequence 9,
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                                                                                                                                          September 21, 2005, 13:41:21; Search time 43 Seconds
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1: /cgn2_6/ptodata/1/iaa/5A_COMB.pep:*

2: /cgn2_6/ptodata/1/iaa/5B_COMB.pep:*

3: /cgn2_6/ptodata/1/iaa/6A_COMB.pep:*

4: /cgn2_6/ptodata/1/iaa/BCTUS_COMB.pep:*

5: /cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*

5: /cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*
                 GenCore version 5.1.6
(c) 1993 - 2005 Compugen Ltd
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US-08-862-607-2
US-08-862-607-2
US-09-203-235-2
US-09-203-235-9
US-09-203-235-9
US-09-203-235-9
US-08-943-136A-4
US-09-588-044-2
US-09-588-044-2
US-09-588-044-2
US-09-588-044-2
US-09-588-044-9
US-09-588-044-9
US-09-588-044-9
US-09-588-048-2
US-09-88-106-493A-2
US-08-106-493A-2
US-08-108-20-264-2
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US-08-821-994-65
US-09-583-110-4712
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                                                                                                                                                                                                                                                                                                                                                                               513545 segs, 74649064 residues
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Gapop 60.0 , Gapext 60.0
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seq length: 200000000
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Match Length
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162
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180
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521
521
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Maximum DB 8
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Sequence 16, Appl
Sequence 16, Appl
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Sequence 16, Appl
Sequence 5, Appli
Sequence 5, Appli
Sequence 9, Appli
Sequence 1267, App
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Sequence 68, Appli
Sequence 5766, Appli
Sequence 8603, Appli
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Sequence 8603, Appli
Sequence 5766, Appli
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Sequence 2, Application US/08352324A

Patent No. 5633149

GENERAL INFORMATION:
APPLICANT: Hawkins, Phillip R.
APPLICANT: Wilde, Craig G.
APPLICANT: Seilhamer, Jeffrey J.
TITLE OF INVENTION: A NOVEL CHEMOKINE EXPRESSED IN
TITLE OF INVENTION: INFLAMED ADENOID, ITS PRODUCTION AND USES
NUMBER OF SEQUENCES: 9
CORRESPONDENCE ADDRESS:
ADDRESSEE: Incyte Pharmaceuticals, Inc.
STREET: 3174 Porter Drive
CITY: Palo Alto
US-08-484-158B-16
US-08-484-596A-16
US-08-480-150A-16
US-08-458-731-16
US-09-149-223A-16
US-09-1149-476-349
US-09-149-476-349
US-09-402-532-10
US-09-402-532-10
US-09-471-276-1267
US-09-288-143-68
US-09-288-143-68
US-09-248-143-68
US-09-631-976-5766
US-09-631-976-5766
US-09-631-976-5766
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MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
COMPUTER: IBM Compatible
COMPUTER: IBM Compatible
COMPUTER: IBM Compatible
COMPUTER: Diskette
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Luther, Barbara J.
REGISTRATION NUMBER: 33,954
REFERENCE/DOCKET NUMBER: PF-0025 US
TELEFAX: 415-855-055
INFORMATION FOR SEQ ID NO: 2:
CENTRANDENMENS: SEQ ID NO: 2:
LENGTH: 109 amino acids
COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE MINIOR COMPUTER: DISKETE
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Gaps

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MKFISTSLLLMLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC
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                                                                                                                                                                                                                US-08-352-324A-9
; Sequence 9, Application US/08352324A
; Patent No. 5633149
; Patent No. 563149
; GENERAL INFORMATION:
APPLICANT: Guegler, Karl J.
APPLICANT: Hawkins, Phillip R.
APPLICANT: Hawkins, Phillip R.
TITLE OF INVENTION: INFLAMED ADDENOID, ITS PRODUCTION AND USES
INDMER OF SEQUENCES: 9
CORRESPONDENCE ADDRESS:
ADDRESSEE INCYTE PHARMACOULICALS, INC.
STREET: 3174 Porter Drive
CITY: Palo Alto
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       .
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                                                                                                                                                          PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
                                                                                                                                                                                                    PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
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MEDIUM TYPE: Diskette E
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FASESEQ VESTON 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/352,324A
FILING DATE: 07-DEC-1994
CLASSIFICATION: 435
ATTORNEY/AGENT INFORMATION:
NAME: Luther, Barbara J.
REGISTRATION NUMBER: 33,954
REFERENCE/DOCKET NUMBER: 33,954
REFERENCE/DOCKET NUMBER: 31,954
REFERENCE/DOCKET NUMBER: 31,954
REFERENCE/DOCKET NUMBER: 31,954
REFERENCE/SOCKET NUMBER: 31,954
REFERE
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Patent No. 5844084
GENERAL INFORMATION:
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TYPE: amino acid
STRANDEDNESS: single
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          , MOLECULE TYPE: peptide US-08-352-324A-9
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Guegler, Karl J.

APPLICANT:

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1 MKFISTSLLLMLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC
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nawkins, Phillip R.

APPLICANT: Wilde, Craig G.
APPLICANT: Seilhamer, Jeffrey J.
TITLE OF INVENTION: A NOVEL CHEMOKINE EXPRESSED IN
TITLE OF INVENTION: INFLAMED ADENOID, ITS PRODUCTION AND USES
NUMBER OF SEQUENCES: 9
CORRESPONDENCE ADDRESS:
ADDRESSEE: Incyte Pharmaceuticals, Inc.
STREET: 3174 Porter Drive
CITY: Palo Alto
STATE: CA
COUNTRY: US
ZIP-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Sequence 9, Application US/08862607
Patent No. 5844084
GENERAL INFORMATION
APPLICANT: Guegler, Karl J.
APPLICANT: Hawkins, Phillip R.
APPLICANT: Wilde, Craig G.
APPLICANT: Sailhamer, Jeffrey J.
TITLE OF INVENTION: INFLAMED ADENOID, ITS PRODUCTION AND USES
TITLE OF INVENTION: INFLAMED ADENOID, ITS PRODUCTION AND USES
CORRESPONDENCE ADDRESS:
ADDRESSEE: Incyte Pharmaceuticals, Inc.
STREET: 3174 Porter Drive
CITY: Palo Alto
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    61 PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              SOFTWARE: FastSEQ Version 1.5
SOFTWARE: FastSEQ Version 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/862,607
FILING DATE: 23-MAY-1997
CLASSIFICATION APPR.
APPLICATION DATE: 08/352,324
FILING DATE: 07-DEC-1994
ATTORNEY AGENT INFORMATION:
NAME: LUther, Barbara J.
REGISTRATION NUMBER: 33,954
RREFERENCE/DOCKET NUMBER: 33,954
RREFERENCE/DOCKET NUMBER: 33,954
RREFERENCE/DOCKET NUMBER: 31,954
RREFERENCE/DOCKET NUMBER: 31,954
RREFERENCE/DOCKET NUMBER: 31,954
RREFERENCE/DOCKET NUMBER: 31,954
RREFERENCE/DOCKET NUMBER: 31,955
RREFERENCE/DOCKET NUMBER: 31,955
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REFERENCE/DOCKET NUMBER: 31,955
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INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
LENGTH: 109 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Inflamed Adenoid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Diskette
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     TYPE: amino acid
STRANDEDNESS: single
                                                                                                                                                                                                                                                                                                                                                                                                                                                               ZIP: 94304
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskett
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           linear
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LIBRARY: INLL.
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1 MKFISTSLLLMLLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC 60
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; Sequence 9, Application US/09203235
; Patent No. 6071701
; GENERAL INFORMATION:
; APPLICANT: Geogler, Karl J.
APPLICANT: Hawkins, Phillip R.
; APPLICANT: Wilde, Craig G.
; APPLICANT: Seilhamer, Jeffrey J.
; TITLE OF INVENTION: INFLAMED ADENOID, ITS PRODUCTION AND USES
; NUMBER OF SEQUENCES: 9
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Incyte Pharmaceuticals, Inc.
; STREET: 3174 POTTER Drive
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 3; Length 109;
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           FILING DATE: 23-MX-1997
APPLICATION NUMBER: 08/352,324
FILING DATE: 07-DEC-1994
ATTORNEY/AGENT INFORMATION:
NAME: Luther, Barbara J.
REGISTRATION NUMBER: 33,954
REFERENCE/DOCKET NUMBER: PF-0025 US
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION INFORMATION:
TELEPHONE: 415-852-0195
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
LENGTH: 109 amino acids
TYPE: amino acid
TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         REFERENCE/DOCKET NUMBER: PF-0025 US
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   OPERATING SYSTEM: DOS
SOFTWARE: FASTSEQ VETSION 1.5
SUBRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/203,235
08/862,607
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              NAME: Luther, Barbara J. REGISTRATION NUMBER: 33,954
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   FILING DATE:
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/862
FILING DATE: 23-MAY-1997
APPLICATION NUMBER: 08/352
FILING DATE: 07-DEC-1994
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                               LIBRARY: Inflamed Adenoid CLONE: 20293
                                                                                                                                                                                                                                                                                                                                             TOPOLOGY: linear
MOLECULE TYPE: polypeptide
IMMEDIATE SURCE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FASTSEQ Version
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
APPLICATION NUMBER:
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APPLICANT: Hawkins, Phillip R.
APPLICANT: Haukins, Phillip R.
APPLICANT: Wilde, Craig G.
APPLICANT: Seilhamer, Jeffrey J.
TITLE OF INVENTION: A NOVEL CHEMOKINE EXPRESSED IN
TITLE OF INVENTION: INFLAMED ADENOID, ITS PRODUCTION AND USES
TOWNERS OF SEQUENCES:
ADDRESSEE: Incyte Pharmaceuticals, Inc.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      61 PRKEIIVWKKNKSIVCVDPQAEWIQRAMEVLRKRSSSTLPVPVFKRKIP 109
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             100.0%; Score 109; DB 2; Length 1
100.0%; Pred. No.:6e-98;
tive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              STREET: 3174 Porter Drive CITY: Palo Alto CONTWINE.
                                                                                                                                                                                                                                                                                                                                           NAME: Luther, Barbara J.
REGISTRATION NUMBER: 33,954
REFERENCE/DOCKET NUMBER: 33,954
RELECOMMUNICATION INFORMATION:
TELEPHONE: 415-85-055
TELEPA: 415-85-055
TELEPA: 415-85-0195
TELEPA: 109 amino acids
LENGTH: 109 amino acids
TYPE: amino acid
STRANDEDNESS: single
                                                                                                                                         SOFTWARE: FastSEQ Version 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/862,607
FILING DATE: 23-MAY-1997
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/352,324
FILING DATE: 07-DEC-1994
ATTORNEY/AGENT INFORMATION:
NAME: Luther, Barbara J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette
COMPUTER: IBM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSEQ Version 1.5
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/203,235
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Sequence 2, Application US/09203235
Patent No. 6071701
GENERAL INFORMATION:
                                                                                    E: Diskette
IBM Compatible
SYSTEM: DOS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match 100.
Best Local Similarity 100.
Matches 109; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      TOPOLOGY: linear
MOLECULE TYPE: peptide
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CLASSIFICATION:
PRIOR APPLICATION DATA:
                                                         COMPUTER READABLE FORM
                                                                                                  COMPUTER: IBM COM
OPERATING SYSTEM:
                                                                             MEDIUM TYPE:
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               COUNTRY: U
ZIP: 94304
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US-09-203-235-2
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Gaps

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GENERAL INFORMATION:
APPLICANT: LI, HADDONG
APPLICANT: KREIDER, BRENT L.
TITLE OF INVENTION: THERAPEUTIC COMPOSITIONS AND METHODS FOR
TITLE OF INVENTION: INHIBITOR-1 (LAI-1), AND CHEMOKINE BETA-11 (CKBETA-11)
NUMBER OF SEQUENCES: 16
CORRESPONDENCE ADDRESS:
ADDRESSEE: STERNE, KESSLER, GOLDSTEIN & FOX. P.1.1.C
1 MKFISTSLLLMLLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC 60
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SSEE: STERNE, KESSLER, GOLDSTEIN & FOX, P.L.L.C.
F: 1100 NEW YORK AVENUE, SUITE 600
WASHINGTON
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               COMPUTER: REALDE FORM
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC COMPAtible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/943,336A
FILING DATE: 03-0CT-1997
PRIOR APPLICATION NUMBER: US 60/027,769
FILING DATE: 04-0CT-1996
PRIOR APPLICATION NUMBER: US 08/724,871
FILING DATE: 04-0CT-1996
PRIOR APPLICATION NUMBER: US 08/464,401
FILING DATE: 05-UW-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/464,401
FILING DATE: 05-UW-1995
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/460,987
FILING DATE: 05-UW-1995
PRIOR APPLICATION NUMBER: US 08/460,987
FILING DATE: 05-UW-1995
PRIOR APPLICATION NUMBER: US 08/460,987
FILING DATE: 08-FEB-1995
ATTONNEY/AGENT INFORMATION:
NAME: CREPER FRICK
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 109; DB 3;
Pred. No. 6e-98;
; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            1488.0380005
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100.0%; Pre
                                                                                                                                                                                                                                                               Sequence 4, Application US/08943336A
Patent No. 6139832
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        NAME: STEFFE, ERIC K.
REGISTRATION NUMBER: 36,688
REFERENCE/DOCKET NUMBER: 14
TELECOMMUNICATION INFORMATION:
TELEPHONE: (202)371-2600
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             INFORMATION FOR SEQ ID NO: 4:
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amino acid
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Best Local Similarity 100.
Matches 109; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         linear
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                                                                                                                                                                                                                                            US-08-943-336A-4
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                                                                                                                                                                                                                                                                                                                                                                 Length 109;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB 3; Length 109;
                                                                                                                                                                                                                                                                                                                                                                                                                         0; Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/982,493
FILING DATE:
CLASSIFICATION: 435
                                                                                                                                                                                                                                                                                                                                                              DB 3;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   100.0%; Score 109; DB 3; ilarity 100.0%; Pred. No. 6e-98; Conservative 0; Mismatches 0
                                                                                                                                                                                                                                                                                                                                                           ; Score 109; DB 3;
; Pred. No. 6e-98;
0; Mismatches (
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 4, Application US/08982493
Patent No. 6110695
GENERAL INFORMATION:
APPLICANT: Gunn, Michael D
APPLICANT: Williams, Lewis T
APPLICANT: Williams, Lewis T
APPLICANT: Cyster, Jason G
ITLE OF INVENTION: Modulating B Lymphocyte (
ITLE OF INVENTION: Receptor Interactions
NUMBER OF SEQUENCES:
CORRESPONDENCE ADDRESS:
ADDRESSEE: SCIENCE & TECHNOLOGY LAW GROUP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               E: SCIENCE & TECHNOLOGY LAW GROUP
75 DENISE DRIVE
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NAME: OSWAN, RICHARD A
REGISTRATION UNDHER: 36,627
REFERENCE/DOCKET UNDHER:
TELECOMMUNICATION INFORMATION:
TELEPHONE: (650) 343-4341
TELEFAX: (650) 343-4342
INFORMATION FOR SEQ ID NO: 4:
SEQUENCE CHARACTERISTICS:
      LELEPHONE: 415-855-0555
TELEFAX: 415-852-0195
INFORMATION FOR SEQ ID NO: 9:
SEQUENCE CHARACTERISTICS:
LENGTH: 109 amino acids
TYPE: amino acid
STRANDEDNESS: single
TOPOLOGY: linear
MOLECULE TYPE: peptide
US-09-203-235-9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ZIP: 94010
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC COMPATIBLE
                                                                                                                                                                                                                                                                                                                                                              100.0%;
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amino acid
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CITY: HILLSBOROUGH
STATE: CALIFORNIA
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
Matches 109; Conserva
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Similarity
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Best Local Simi
Matches 109;
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1 MKPISTSLLLMLLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC
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                                                                                                                                                             APPLICANT: Guegler, Karl J.
Hawkins, Phillip R.
Wilde, Craig Ge
Seilhamer, Jeffrey J.
TITLE OF INVENTION: A NOVEL CHEMOKINE EXPRESSED IN
INFLAMED ADENOID, ITS PRODUCTION AND USES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ö
PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
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                                                                                                                                                                                                                                                                                                                          CORRESPONDENCE ADDRESS:
ADDRESSEE: Incyte Pharmaceuticals, Inc.
STREET: 3174 Porter Drive
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             CURRENT APPLICATION DATA:

APPLICATION NUMBER: U5/09/588,044
FILING DATE: 05-Jun-2000
CLASSIFICATION: «UNKNOWN»
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/352,324
FILING DATE: 07-DEC-1994
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         CLONE: 20293
SEQUENCE DESCRIPTION: SEQ ID NO: 2:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MEDIUM TYPE: Diskette
COMPUTER: 1BM Compatible
OPERATING SYSTEM: DOS
SOFTWARE: FastSEQ Version 1.5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ATTORNEY/AGENT INFORMATION:
NAME: Luther, Barbara J.
REGISTRATION NUMBER: 33,954
REFERENCE/DOCKET NUMBER: PF
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          LIBRARY: Inflamed Adenoid
                                                                                                       Sequence 2, Application US/09588044
Patent No. 6692920
GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         LENGTH: 109 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Sequence 9, Application US/09588044
Patent No. 6692920
GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           APPLICANT: Guegler, Karl J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        TOPOLOGY: linear MOLECULE TYPE: polypeptide
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        TELEFAX: 415-852-0195
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  STRANDEDNESS: single
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               COMPUTER READABLE FORM
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Matches 109; Conservative
                                                                                                                                                                                                                                                                                                    NUMBER OF SEQUENCES:
                                                                                                                                                                                                                                                                                                                                                                                         CITY: Palo Alto
                                                                                                                                                                                                                                                                                                                                                                                                                                      SD
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                                                                                                                                                                                                                                                                                                                                                                                                                                                        ZIP: 94304
                                                                                                                                                                                                                                                                                                                                                                                                                                      COUNTRY:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      RESULT 11
US-09-588-044-9
                                                                                     US-09-588-044-2
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                                                                                                                                                                                                                                APPLICANT: LI, HAODONG
KREIDER, BRENT L.
TITLE OF INVENTION: THERAPEUTIC COMPOSITIONS AND METHODS FOR
TITLE OF INVENTION: THERAPEUTIC COMPOSITIONS AND METHODS FOR
TREATING DISEASE STATES WITH LEUKOCYTE ADHESION
INHIBITOR-1 (LAI-1), AND CHEMOKINE BETA-11 (CREETA-11)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ö
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                      PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
                                               61 PRKEIIVWKKNKSIVCVDPQAEWIQRAMEVLRKRSSSTLPVPVFKRKIP 109
                                                                                                                                                                                                                                                                                                                                            NUMBER OF SEQUENCES: 16
CORRESPONDENCE ADDRESS:
ADDRESSE: STERNE, KESSLER, GOLDSTEIN & FOX, P.L.L.C.
STREET: 1100 NEW YORK AVENUE, SUITE 600
CITY: WASHINGTON
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      PRKEIIVWKKNKSIVCVDPQAEWIQRAMEVLRKRSSSTLPVPVFKRKIP 109
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Best Local Similarity 100.0%; Pred. No. 6e-98;
Matches 109; Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ATTORNEY AGENT INFORMATION:
NAME: STEFFE, ERIC K.
REGISTRATION UNDRER: 36,688
REFERENCE/DOCKET NUMBER: 1488.0380006
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             APPLICATION NUMBER: US 08/943,336
FILING DATE: 03-0CT-1997
APPLICATION NUMBER: US 60/027,769
FILING DATE: 04-0CT-1996
APPLICATION NUMBER: US 08/724,871
FILING DATE: 04-0CT-1996
APPLICATION NUMBER: PCT/US96/09572
FILING DATE: 05-JUN-1996
APPLICATION NUMBER: US 08/464,401
FILING DATE: 05-JUN-1996
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       COMPUTER: Floppy disk
COMPUTER: IBM PC compatible
COMPUTING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Ver
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/635,899
FRIOR APPLICATION DATA:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              FILING DATE: 05-JUN-1995
APPLICATION NUMBER: PCT/US95/01780
FILING DATE: 08-FEB-1995
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ID NO: 4:
                                                                                                                                                                    Sequence 4, Application US/09635899
Patent No. 6485719
GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            LENGTH: 109 amino acids TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TOPOLOGY: linear
MOLECULE TYPE: protein
SEQUENCE DESCRIPTION: SEQ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 INFORMATION FOR SEQ ID NO: 4: SEQUENCE CHARACTERISTICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ZIP: 20005
COMPUTER READABLE FORM:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                        STATE: DC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               COUNTRY:
                                                                                                                                                  US-09-635-899-4
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NAME: LUTHER, BARBARA J.
REGISTRATION NUMBER: 33954
REPERENCE, DOCKET NUMBER: PI
TELECOMMUNICATION INFORMATION
TELEPHONE: 415-855-0555
SYSTEM: MS-DOS
WORD PERFECT 5.1
                                                                                                                                                                                                                                                                                                                                                                           INFORMATION FOR SEQ ID NO: 4: SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                         LENGTH: 109 AMINO ACIDS TYPE: AMINO ACID STRANDEDNESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 TELEFAX: 415-852-0195
INFORMATION FOR SEQ ID NO:
                                                                                                                                        PRIOR APPLICATION DATA:
APPLICATION NUMBER:
FILING DATE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TOPOLOGY: LINEAR MOLECULE TYPE: PROTEIN
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APPLICANT: LI. ET AL.
APPLICANT: LI. ET AL.
TITLE OF INVENTION: Human Chemokine Beta-11 and Human Chemokine Alpha-1
NUMBER OF SEQUENCES: 16
CORRESPONDENCE ADDRESS:
ADDRESSEE: CARELLA, BYNNE, BAIN, GILFILLAN,
ADDRESSEE: CECCHI, STEWART & OLSTEIN
STREET: 6 BECKER FARM ROAD
CITY: ROSELAND
STATE: NEW JERSEY
COUNTRY: USA
ZIP: 07068
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1 MKFISTSLLLMLLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC 60
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Hawkins, Phillip R.
Wilde, Craig G.
Seilhamer, Jeffrey J.
TITLE OF INVENTION: A NOVEL CHEMOKINE EXPRESSED IN
INFLAMED ADENOID, ITS PRODUCTION AND USES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
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100.0%; Score 109; DB 4; Length 10
Best Local Similarity 100.0%; Pred. No. 6e-98;
Matches 109; Conservative 0; Mismatches 0; Indels
                                                                                                                                           CORRESPONDENCE ADDRESS:
ADDRESSEE: Incyte Pharmaceuticals, Inc.
STREET: 3174 Porter Drive
CITY: Palo Alto
                                                                                                                                                                                                                                                                                                                        COMPUTER: Diskette

COMPUTER: IBM Compatible

OPERATING SYSTEM: DOS

SOFTWARE: FastSEQ Version 1.5

CURRENT APPLICATION DATA:

APPLICATION UNBER: US/09/588,044

FILING DATE: 05-Jun-2000

CLASSIPICATION: <Unknown>
PRIOR APPLICATION NUMBER: 08/352,324

FILING DATE: 07-DEC-1994

ATTORNEY/AGENT INFORMATION:

NAME: Luther, Barbara J.

REFERRATION NUMBER: 33,954

REFERRATION NUMBER: 13,954
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       MOLECULE TYPE: peptide
SEQUENCE DESCRIPTION: SEQ ID NO: 9:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       TELECOMMUNICATION INFORMATION:
TELEBHONNE: 415-855-0555
TELEBAX: 415-852-0195
INFORMATION FOR SEQ ID NO: 9:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SEQUENCE CHARACTERISTICS:
LENGTH: 109 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TYPE: amino acid
STRANDEDNESS: single
                                                                                                                                                                                                                                                                                                            COMPUTER READABLE FORM:
                                                                                                                       NUMBER OF SEQUENCES: 9
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MEDIUM TYPE: 3.5 INC!
COMPUTER: IBM PS/2
                                                                                                                                                                                                                                                             COUNTRY: US
                                                                                                                                                                                                                                                                                 ZIP: 94304
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1 MKFISTSLLIALLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC 60
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TITLE OF INVENTION: A NOVEL CHEMOKINE EXPRESSED IN INFLAMED
TITLE OF INVENTION: ADENOID, ITS PRODUCTION AND USES
NUMBER OF SEQUENCES: 9
CORRESPONDENCE ADDRESS:
ADDRESSEE: INCYTE PHARMACEUTICALS, INC.
STREET: 3174 Porter Drive
CITY: Palo Alto
STATE: CA
COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 5; Length 109;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match 100.0%; Score 109; DB 5; : Best Local Similarity 100.0%; Pred. No. 6e-98; Matches 109; Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           COMPUTER READBLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: WordPefect 6.1/MS-DOS 6.2
CURRENT APPLICATION DATA:
FLING DATE: 07-DEC-1995
                                                                                                                                                                                                                                                                               325800-272
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US95/01780
FILING DATE: Concurrently
CLASSIFICATION:
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CLASS/FICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/352,324
FILING DATE: 07-DEC-1994
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Sequence 2, Application PC/TUS9516144 GENERAL INFORMATION:
                                                                                                                                                                                         ATTORNEY/AGENT INFORMATION:
NAME: FERRANCY, GREGORY D.
REGISTRATION NUMBER: 36,134
REFERENCE/DOCKET NUMBER: 325;
TELECOMUNICATION INFORMATION:
TELECHONE: 201-994-1700
TELEFAX: 201-994-1744
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Search completed: September 21, 2005, 13:50:03 Job time : 44 secs
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             APPLICANT: INCTE PHARMACEUTICALS, INC.
TITLE OF INVENTION: A NOVEL CHEMOKINE EXPRESSED IN INFLAMED
TITLE OF INVENTION: A DENOID, ITS PRODUCTION AND USES
TOWNER OF SEQUENCES: 9
CORRESPONDENCE ADDRESS:
ADDRESSES: INCYTE PHARMACEUTICALS, INC.
STREET: 3174 Porter Drive
                                                                                                                                                                                                                                      DB 5; Length 109;
                                                                                                                                                                                                                                                                                Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match 100.0%; Score 109; DB 5; Sect Local Similarity 100.0%; Pred. No. 6e-98; Matches 109; Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: WordPefect 6.1/MS-DOS 6.2
CURRENT APPLICATION DATA:
APPLICATION NUMBER: PCT/US95/16144
FILING DATE: 07-DEC-1995
CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/352,324
FILING DATE: 07-DEC-1994
ATTORNEY/AGENT INFORMATION:
NAME: LUTHER, BARBARA J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ;.Sequence 9, Application PC/TUS9516144; GENERAL INFORMATION:
LENGTH: 109 amino acide
TYPE: amino acide
TYPE: amino acide
STRANDEDNESS: single
TYPEOLOGY: linear
MOLECULE TYPE: polypeptide
IMMEDIATE SOURCE:
LIBRARY: Inflamed Adenoid
CLORRARY: 10293
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 TELECOMMUNICATION INFORMATION:
TELEPHONE: 415-855-0555
TELEFAX: 415-852-0195
INFORMATION FOR SEQ ID NO: 9:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       NAME: LUTHER, BARBARA J. REGISTRATION NUMBER: 33954
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 REFERENCE/DOCKET NUMBER:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             LENGTH: 109 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ZIP: 94304
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   amino acid
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                                                                                                                                                                                         PCT-US95-16144-2
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STATE: CA
COUNTRY:
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1 MKFISTSLLLAMLVSSLSPVQGVLEVYYSLRCRCVQESSVFIPRRFIDRIQILPRGNGC 60

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Sequence 7722, Application US/09949016

Fatent No. 6812339

GENERAL INFORMATION:

APPLICANT: VENTER, J. Craig et al.

TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF

TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF

TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES THEREOF

CURRENT APPLICATION NUMBER: US/09/949,016

CURRENT FILING DATE: 2000-04-14

FRIOR PAPLICATION NUMBER: 60/231,768

FRIOR FILING DATE: 2000-10-03

FRIOR FILING DATE: 2000-10-03

FRIOR PILING DATE: 2000-09-08

NUMBER OF SEQ ID NOS: 207012

SOFTWARE: FRAELSEQ for Windows Version 4.0
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1 MKFISTSLLLMLLVSSLSPVQGVLEVYYTSLRCRCVQESSVFIPRRFIDRIQILPRGNGC 60
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                                                                               PRKEIIVWKKNKSIVCVDPQAEWIQRMMEVLRKRSSSTLPVPVFKRKIP 109
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               100.0%; Score 109; DB 4; Length 115; 100.0%; Pred. No. 6.3e-98; ive 0; Mismatches 0; Indels (
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity 100.
Matches 109; Conservative
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ORGANISM: Human
                                                                                                                                                                                                             US-09-949-016-7722
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LENGTH: 115
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